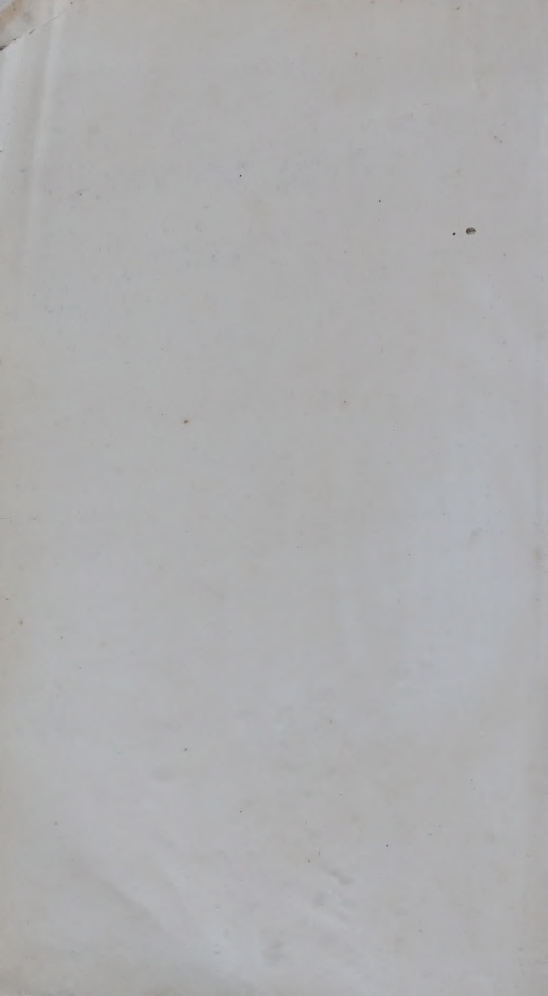


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REINHOLD ORGANIC CHEMISTRY AND BIOCHEMISTRY TEXTBOOK SERIES

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FOOD CHEMISTRY

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CALVIN A. VANDERWERF

Consulting Editor

FOOD CHEMISTRY

LILLIAN HOAGLAND MEYER

Professor and Head, Department of Chemistry Western Michigan University Kalamazoo, Michigan



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Preface

THE RAPID GROWTH of the food industry into big business and the changes in the number of items on the grocers' shelves, the many ready-to-eat products, the new controls on food additives, and the attempts to standardize some food articles all serve to emphasize the growing importance of the chemistry of foods. For centuries the production of food followed traditional procedures, and new ideas or methods were largely the result of empirical trial and error or accidental discoveries. Now a complete revolution in growing, processing, and marketing of food is here. The accumulation of scientific knowledge about food composition has been, in the main, slow. Dependent as the science of food is on biology, bacteriology, and mycology as well as chemistry, it was necessary for all sciences to develop to the point where elucidation of the complex mixtures encountered in foods became accurate and meaningful. Fortunately, research of significance to food chemistry is now appearing not only in journals devoted exclusively to food problems but also in those in the fields of biology, chemistry, chemical engineering, and even physics.

It is the author's belief that an attempt to consolidate the fundamentals of food chemistry with recent advances in the food industry is now appropriate. Although there have been splendid books available for many years in the area of food analysis and more lately in food technology, we have not had one that unifies these areas with basic chemistry in a simple fashion. The present book is designed to meet this need. It is intended primarily as a text in food chemistry for undergraduate students in home economics, food technology, and chemistry. However it should be useful also as a reference book for students and research workers in these fields.

Thus the aim of the book is to provide a unified picture of foods from a chemical standpoint. The primary emphasis is on the composition of foods

and the changes that occur when they are subjected to processing. author has tried to keep in mind undergraduates who have had a cour organic chemistry; however, an effort has been made to provide neces background chemistry and sufficient explanations for students to pro without additional references. Some brief descriptive orientation is g on the structure of plants and animals, but this is kept quite simple ar used to promote greater unity of presentation. Occasionally, materia food habits is included to add interest and to round out the close relat ship of food chemistry to everyday life. The chapter on food additives the appendix on permissible food additives are special features of this that provide up-to-date information on this vital topic. At the end of echapter are bibliographical references to aid the reader who wishes to vestigate further the respective areas of food chemistry.

Thanks are due to many people for the successful completion of book. Calvin A. VanderWerf and Betty Watts were especially helpfu their critical evaluation of the manuscript and Betty Taylor and Cu

Meyer for their suggestions, criticisms, and continued interest.

LILLIAN HOAGLAND ME

Kalamazoo, Michigan June, 1960

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Development of Food Chemistry

HE ORIGINS OF SCIENCE go back to prehistory when man could not write a ecord of his successes and his failures, his trials and experiments, and his aditions. We assume that as man emerged as a thinking animal he began explore the world around him, to wonder at its operation, and finally to y to control it. Consequently, it seems that as the centuries roll by nan's understanding and control of his environment, although still incomlete, continue to improve. In the early days every science had a tenuous art—even one as dependent on the development of mathematics and labratory procedures as chemistry. At this time man had learned only a few nemical skills, some of which were in food chemistry. Probably they ere accidently discovered, then carefully passed on from one generation to nother so that by the time writing was invented man could not only make ottery and soap, cook food, and smelt a few metals but also make wine nd vinegar. These processes were hedged around with superstitions, but e vintner could control the process; although he had never heard of a east or an Acetobacter, he knew how to handle the fermenting fruit.

In the seventeenth century, as laboratory procedures and an appreciaon of the scientific method developed and as scientists began to exnange information, chemistry began to emerge as a true science. A condetable accumulation of information and a beginning study of foods offected the general interest in the composition of the entire world. Some the early experiments attempted to separate food components, to study be pigments that give color to the world, or to find the "life-giving" impound. As time passed these studies were successful to a limited extent. If the eighteenth century simple compounds were isolated, and during the est half of the nineteenth century the old idea that the living and the onliving world were completely different was abandoned. The terms "inorganic" and "organic" reflect this early belief in two types of chemistry. When some compounds such as sodium chloride were found in the "inorganic" earth as well as in the blood and urine of a living animal and when "organic" compounds were shown to obey the same laws as "inorganic," the idea was discarded. However, a complete understanding of the composition and the changes in composition of such complex mixtures as living cells was impossible at this stage.

The nineteenth century saw the development of organic chemistry, analytical chemistry and physical chemistry—all essential to the growth of food chemistry and our understanding of it. The pace at which discoveries were made and at which advances occurred increased throughout the century. The field of carbohydrates began to fit together, proteins were recognized, and many other compounds of importance in food chemistry were studied.

Biology also saw rapid growth during the nineteenth century; understanding of this field was also required before much progress could be made in food chemistry. The Cell Theory and the Theory of Evolution were fundamental to much insight into either animal or plant materials. These theories and the bulk of knowledge supporting them gave scientists an appreciation of the complex nature of cells as well as the common patterns that cells may have through their common heritage and the different patterns that separate families, species, and even varieties.

In the twentieth century, the passage of the Pure Food and Drug Act by the United States Congress in 1906 had a great influence on the channels that research in food chemistry took. Harvey Wiley, shocked at the food which could be legally sold in the United States, campaigned for years for the passage of a bill to restrict filth, decay, and adulteration in food. After the passage of the bill, there came a long fight for enforcement. As always in the United States, a company or individual must be proved guilty; consequently under the Pure Food and Drug Act proving that a food did not come up to standard was necessary. The act was originally administered by the Department of Agriculture; chemists in this department worked out many tests to detect adulteration and to set up standards. Eventually the methods developed for foods and other products were published as the Official Methods of Analysis of the Association of Official Agricultural Chemists. The eighth edition, published in 1955, runs to 1,008 pages.

A phenomenal growth in our knowledge of biochemistry has occurred in recent years. Metabolic pathways in plants and animals are being clarified, the structure of large molecules is yielding to study, applications of physical chemical principles become easier and more meaningful every day. These

developments result in a better understanding of the biological materials which we use as food. It is not yet possible to separate completely all of the compounds which make up a cell or to describe even sketchily all of the reactions that occur in a cell, but the body of knowledge has now become large and is rapidly increasing. Probably only a rather small per cent of the reactions that occur when a food is cooked or processed in some way are known, but more are being studied every day.

Through the centuries that man has known fire, cooking procedures have developed empirically. Many of these processes were the result of trial and error, while some must have been discovered accidentally. When a man stumbled on a process, cheese making for example, he handed it down to his descendants. Sometimes it underwent modification, but often the same rules have been followed for generation after generation. These procedures have seldom been the result of scientific experimentation where a series of controlled batches are used and where variations are planned and at least partially understood. Empirical processes usually contain a number of procedures which are essential for success, but sometimes there are procedures which have little or no bearing on the problem. The salt added in salt-rising bread was thought for generations to be important in keeping the "starter" fresh and sweet. Actually, it had little effect on the bacteria and yeasts present.

The food industry developed from small operations, sometimes built around one small kitchen with one stove, or a little butcher or grocery shop. Many industrial processes also developed by trial and error methods. Today, with nationwide industries and with the growth of a consumer market which expects and demands a uniform product, it is necessary to control processes carefully. This control demands understanding of the process. Some industries have been very active in attempting to develop a scientific basis for their procedures. There has been, for example, a number of studies on orange juice concentration. While these researches are directed to solving a problem of the orange juice industry, they have contributed to our knowledge of some basic problems in food chemistry. Browning of orange juice during processing is one of the problems which plagues the processors; the solution of this problem may have a bearing on browning reactions in other foods.

INDIVIDUAL VARIABILITY

In biochemical research it has been recognized for years that an adequate sample must include a fairly large number of individual subjects since variability between members of the same plant or animal species is com-

mon. Thus we can expect every tomato on a single vine to contain ascoacid, but we cannot expect every one to have the identical concentrator of ascorbic acid. Added to this is the variability which is introduced variety and growing conditions. In animals we also find a range of variety and exactly the same composition. When we analyze a composition we find that each sample has exactly the same composition; but when analyze a plant or animal, we find that the values fall into a range.³

An appreciation of variability is essential for anyone concerned variability is essential for anyone concerned variable food chemistry, since it requires that an adequate sample include a number of individuals and also that samples be adequately mixed. If broce for instance, is analyzed, a number of samples from different plants, gree in different locations to the same degree of maturity should be used. The buds and stem must be chopped finely, well mixed, and representate samples withdrawn. The importance of adequate sampling cannot be overwhasized. Too often conclusions have been drawn from small samp These conclusions are often not valid.

INDIVIDUAL UNIFORMITY

On the other hand, both plant and animal tissues have many things common. Since all living things appear to have sprung from the san though distant, ancestor, they possess some compounds and some met bolic pathways in common. With mutations and the occurrence of var tions or even the development of new species, some differences in che istry must necessarily be present. Often the chemical variation is signicant but relatively small. Thus every living cell contains a number proteins. These proteins are not identical, but they have many propert and some aspects of their structure in common. Another example of ur formity is the metabolic pathway of glucose or glycogen to pyruvic ac This series of reactions is used by all plant and animal cells which habeen studied thus far.

An appreciation of uniformity is also important in food chemistry sin the accumulated knowledge is still scanty. We often infer that if a reation explains an observation in one food, it will apply to another. The the study of the browning of orange juice may tell us something about other browning reactions. It is, however, necessary to maintain a critical as well as a skeptical attitude and to recognize that although transfer information from one food to another may be valid, it must be verified.

METHODS OF SAMPLING

When samples of foods are taken for analysis, they are usually fra

mented, minced, or macerated in some manner. Dry samples are sometimes ground in a ball mill, sometimes by some other device; wet samples such as plant and animal tissue are usually ground or macerated. Loss of juice through the squeezing of the grinder knives must be avoided or the juice squeezed out must be returned and mixed with the sample. It is essential that adequate mixing of the sample occur so that the composition does not vary from one part to another. With liquids and pulverized materials, it is fairly easy to pour the sample from one container to another or to mix by stirring. With solid material such as a frozen food, it is necessary to take drillings or chucks from different parts of the package, and after proper treatment such as thawing, mix them thoroughly. When the package or batch is exceedingly large, such as a boxcar, it is necessary to take a number of samplings. Various devices have been developed for taking cores of samples at different places in the batch, which are then carefully mixed and representative samples removed. Often, quartering is used. The sample is spread on a sheet of paper in a cone and flattened to a circle. If the sample is very large, a blanket or sheet must be used. The circle is divided into quarters and two opposite segments, quarters 1 and 3, removed. These in turn are quartered, but segments 2 and 4 are saved this time. The process is repeated until the sample is reduced to a size suitable for analysis. The sample is mixed and often pulverized further between each quartering.

MOISTURE IN FOODS

The most abundant compound, and the one which is almost always present in foods, is water. Occasionally a food such as an oil will be dry; but even crystallized substances which are relatively pure, such as sugar and salt, contain small amounts of water adsorbed on the surfaces of the crystals. Cellular material, whether plant or animal, contains an abundance of water. In leafy green vegetables there is 90 or more per cent water, while even in cooked meat where some water has been driven off the amount is between 50 and 65 per cent.

In plants and animals water is present in the circulating fluid—the saper the blood and lymph—between the cells as intercellular fluid and within the cells. If an animal is cut, fluid drips out; even when the stalk of a plant is cut, fluid can sometimes be seen to drip or ooze out of the vessels which carry the sap. Within the cells water is plentiful in the cytoplasm. In some types of plant cells special cell sap vacuoles contain solutions of compounds dissolved in water.

The water which is present in foods may be held (1) as free liquid in which substances are dissolved or dispersed, (2) as hydrates. (3) as imbibed

water in gels, or (4) by adsorption on the surfaces of solids. Examples of the first type are found in cytoplasm, intercellular fluid, and any of the circulating fluids of tissues. In the second type, hydrates form either when hydrogen bonds are established between water molecules and ions or molecules which contain oxygen or nitrogen or when the unshared electrons of the oxygen are coordinated with an ion. Starches, proteins, and many other organic compounds important in foods, as well as salts, form hydrates. Imbibed water may not be different from water held as a hydrate. Some substances pick up water and swell when they come in contact with water. They are said to "imbibe" water; this may be accomplished by hydrogen bonding. The fourth type of water is held on all surfaces exposed to air in which water vapor is present. "Dry" cocoa holds water and air on the surface of the particles. Solids which are very finely divided have a very large surface area and consequently have a high adsorptive capacity.

Hydrogen Bonding

Water molecules are able to attach themselves to other molecules by means of a hydrogen bond. Water molecules are dipoles in which the hydrogen atoms are slightly positive and the oxygen atom slightly negative. Within a water molecule the hydrogen atoms are bonded to the oxygen by a covalent pair of electrons, but the angle between these atoms is 105°. Thus the four unshared electrons of the oxygen atom are on one side and create a slight negativity. The oxygen atom holds the electrons closely and draws them slightly away from the proton. As a result, the oxygen is slightly negative in comparison to the hydrogens which are positive.

The hydrogen bond or bridge is able to form whenever a slightly positive hydrogen approaches an atom such as oxygen which tends to be slightly negative. The small size of the hydrogen atom allows it to come very close to this atom and establish a weak bond or bridge to it.

In pure water two molecules associate at some temperatures when one hydrogen is attracted to the slightly negative oxygen of another mole-

cule of water and association occurs through the hydrogen bond. The strength of the bond between the two molecules is not nearly as great as the strength of the covalent bonds holding the two hydrogens and oxygen together, but it is much greater than the van der Waal's forces which hold molecules close together in liquids and even closer in solids.

$$H \sim O - H = O$$

Energy in the form of heat must be supplied in order to disrupt hydrogen bonds. The relatively high boiling point of water, a very low molecular weight compound, is explained on the basis of the heat required to break hydrogen bonds.

Not only can hydrogen bonding hold one water molecule to another, it can also cause association or hydrate formation between water and compounds which have polar oxygens. A compound such as methanol, CH₃OH, or a carbohydrate which has a hydroxyl group will bind water through hydrogen bonding. Indeed, pure methanol will associate with itself through hydrogen bonding. In large complex molecules which are coiled and folded water will penetrate within the molecule, forming hydrates through hydrogen bonding and causing a change in the size and shape of the molecule.

The nitrogen atom is frequently linked to another nitrogen or an oxygen atom through a hydrogen bond. Like oxygen, nitrogen tends to be slightly negative, and it will have an attraction for the slightly positive hydrogen of a molecule such as water and bond it to the nitrogen. An amino group in a compound such as a protein will associate with the hydrogen of a hydroxyl group in, say, methanol. The strength of the hydrogen bond is not as strong in this case as it is with oxygen since nitrogen is not as negative as oxygen and the hydrogen is not held as closely.

Hydrates form with those metallic ions which tend to form complexes. The unshared electrons of the oxygen atom will fill out the shells of the

ion and hold water to it. Sodium, magnesium, calcium, and many other ments exist as hydrated ions in solution. The hydration of the hydroion has been emphasized in recent years by use of a special name, "hydronium ion."

Bound Water

Many years ago it was discovered that some of the water in cellular terial can be readily removed by pressing or heating, while some can Gortner attempted to discover why pine needles did not freeze in the verter. He found that needles gathered at that time of year have little was which can be squeezed out, although in the summer needles this is not to In summer less of the water is "bound." In all cellular material ther some bound water.

In working with water in food it is important to remember that some the water may be bound and very difficult to separate. Often it is extrem difficult to separate the water without decomposing other molecules pres in the sample.

Determination of Moisture

In some foods the determination of moisture is relatively simple. It sample is weighed and heated in an oven to constant weight. The different in weight is the water which has evaporated. The sample is usually weight into a flat bottomed, shallow dish made of aluminum or similar mate which will not react with the food nor pick up water readily. The or must be thermostatically controlled and is usually set at 100°C or 105°A thin layer of sand, pumice, or asbestos is often added to the bottom of the dish to support the food particles and accelerate drying.

Many foods decompose to some degree if they are heated to 100°C. T is true, for example, of all foods which contain fructose. It is necessate to dry them in a vacuum oven where the temperature is maintained a lower figure and the pressure is reduced to facilitate loss of moisture. Alternative method with foods sensitive to heat is use of a vacuum descator with sulfuric acid as the drying agent. The samples are again drive constant weight.

Those foods which contain volatile compounds other than water must treated by another method. None of the weight-loss methods are adequated to differentiate between loss of water and loss of some other volatile sustance. The immiscible solvent distillation method can be used for the purpose. The sample is placed in a flask which is connected with a reflect condenser equipped with a distillate trap. (See Figure 1.1.) The sample covered with a suitable solvent and the trap filled with the solvent. The sample covered with a suitable solvent and the trap filled with the solvent.

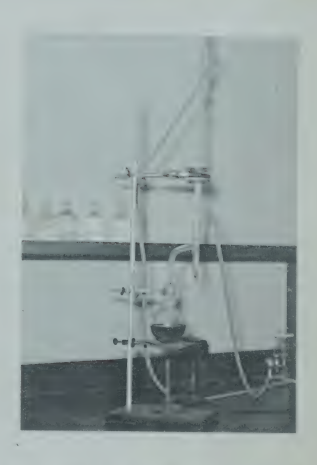


FIGURE 1.1 APPARATUS FOR THE DETERMINATION OF MOISTURE IN FOODS. An immiscible liquid such as toluene boils off with the water. Water is trapped and measured in side vessel.

solvent must be immiscible with water so that as they distill separation of the two liquids can occur. Toluene is most commonly used, although xylene and heptane are sometimes employed. The flask is heated and the vapors of water and solvent are condensed by the condenser and drop into the trap. The lighter solvent flows over into the flask, but the water is captured. If the trap is calibrated, the amount of water distilled out of the sample can be read directly.^{1,2}

Nuclear magnetic resonance has been developed into a rapid and simple method for the determination of moisture in samples, particularly solids. The equipment is rather expensive, but it is possible for an untrained person to make numerous determinations in a very short time. The instrument can be calibrated to read per cent moisture directly. Nuclear-magnetic-conance measurements depend on the magnetic behavior of the nuclei of atoms. All nuclei are positively charged owing to their load of protons and many spin either clockwise or counterclockwise. The rotation of a charged body creates a magnetic field. Only those nuclei which have an even number of protons and neutrons (${}_{6}C^{12}$, ${}_{8}O^{16}$, and ${}_{16}S^{32}$) do not appear to have an angular momentum or create this tiny magnetic field. If spinning nuclei are placed in the field of a magnet, the magnetic field exerts a torque on hem and tends to align them with the field. They absorb radio-frequency

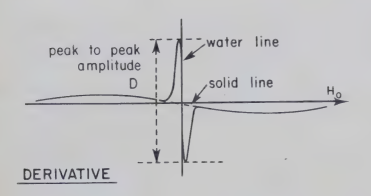


FIGURE 1.2 NUCLEAR MAGNETIC RESONANCE DERIVATIVE OF ABSORPTION CURVE OF MATERIAL CONTAINING SOLID AND WATER. The contribution of protons in the solid ("solid line") is small compared to those in water ("water line").

Courtesy of H. Rubin,
Ridgefield Instrument Group,
Schlumberger Corp.

energy and precess at a definite frequency like tiny gyroscopes. The frequency of the energy which can be absorbed is characteristic of each isotope. Thus in a magnetic field of 1700 gauss, hydrogen will absorb energy at 7.25 megacycles and change to another magnetic energy level. The amount of energy absorbed will be proportional to the quantity of hydrogen present in the sample.

The apparatus is designed to measure the absorbence of energy when the sample is placed inside a coil supplied with radio-frequency current. The coil is mounted between a large permanent magnet. The field strength of the magnet is slowly varied; and when it crosses the NMR value for the

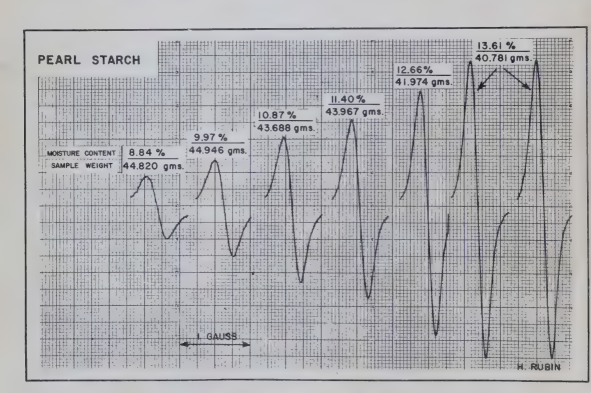


FIGURE 1.3. NUCLEAR MAGNETIC RESONANCE. Curves of moisture in pearl starch.

Courtesy of H. Rubin, Ridgefield Instrument Group, Schlumberger Corp.

particular isotope, the nuclei absorb energy and shift magnetic energy levels. The amount of energy absorbed is recorded.

Many compounds other than water contain hydrogen, but the nature of the compounds and particularly the physical state of the compounds have an effect on the width of the line and shape of the curve produced. In liquids the motion of molecules is so rapid that the effect of the magnetic field created by neighboring nuclei is smoothed out. In many samples water is the only liquid present.

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CHAPTER TWO

Fats and Other Lipids

LIPIDS ARE ONE OF THE LARGE GROUPS of organic compounds which are of great importance in the food we eat because they are readily digested and utilized in the body. They are widely distributed and almost every natural food has considerable quantities of them. Fruits and vegetables are not ordinarily thought of as sources of lipids, but we find that most of them contain between 0.1 and 1 per cent total lipid, usually listed as fat.

A few fruits and vegetables are rich sources of these compounds. Thus an avocado contains an average of 20 per cent lipid while a ripe olive has about 19 per cent. Whole-grain cereals have from 1 per cent for whole barley on the dry basis to 7.4 per cent for dry oatmeal, while nuts are very rich in lipid. Pecans average 73 per cent and English walnuts about 64 per cent. But the natural foods which contribute the largest amounts of these compounds to our diet are the animal products—meats and fowl, milk and milk products, and eggs.

Fats are also often added to foods during their preparation, either as shortening, as a method of transferring heat in frying, or for flavor and richness in preparing vegetables, puddings, etc. Occasionally lecithin is added as an emulsifying agent, but the fats are by far the most important additives among the lipids.

OCCURRENCE IN FOODS AND COMPOSITION

The American diet is unusually rich in fats and other lipids. During recent years the percentage of calories derived from fat has increased markedly in the average diet. Many nutritionists and physicians are alarmed at this high consumption and consider it unwise. Table 2.1 shows the consumption of fats in the United States in recent years. The problem of diet fat is briefly discussed on page 62.

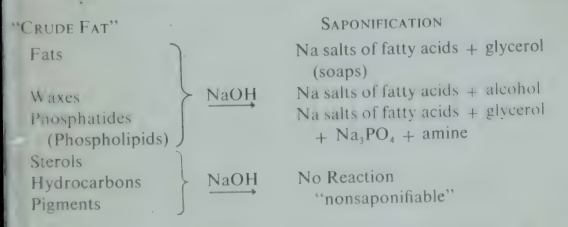
The lipid fraction of a food is usually separated from the other compounds present in the food by extraction with a solvent such as petroleum ether, ethyl ether, chloroform, or benzene, and is reported either as "ether-soluble fraction" or "crude fat." Actually the fraction contains not only true fats but also waxes, complex lipids such as phospholipids, derived lipids such as the sterols, and many pigments, hormones, and volatile oils.

TABLE 2.1. CONSUMPTION OF FOOD FATS AND OILS, EXCEPT BUTTER*

	1955-56	1956-57
	(Million Lb)	
Stocks, Beginning of Period	\944	949
Produced	8347	8209
Total	, 9291	9158
Stocks, End of Period	949	776
Total Disappearance	8342	8382
Export	2002	1937
Domestic Use	6340	6445

^{*}From J. Am. Oil Chemists' Soc., Nov., 1957, p. 8.

Occasionally this crude fat fraction is further separated. The sample is boiled with an alkali such as sodium hydroxide and by saponification the fats, waxes, compound lipids, and free fatty acids then form soaps. These disperse in the water layer and other products such as glycerol and phosphates dissolve in water, but the sterols, pigments, hydrocarbons, etc. are insoluble. The "saponifiable fraction" thus includes all lipids except the sterols, pigments, and hydrocarbons. Saponification of the "saponifiable fraction" (fats, waxes, and phosphatides) is shown diagrammatically below:



I've fat content reported in foods is often this "crude fat" and represents total lipid content rather than true fat.

Little information on the distribution of lipids in the crude fat fraction of most foods is generally available. In animal tissues true fat is found primarily in the adipose tissues, while the active tissues, those which use considerable oxygen and produce much carbon dioxide—the muscles, nervous tissue, and glands—contain relatively little fat in the lipid fraction and much more of the complex lipids and sterols.

The fats and oils of commerce are for the most part mixtures of fats, the triglycerides of fatty acids.

$$C_{17}H_{35}COOCH_2$$
 $C_{17}H_{31}COOCH$
 $C_{15}H_{31}COOCH_2$

A Fat

Some of the purified oils and fats contain only very small amounts of other compounds, and these, for the most part, have little or no effect on flavor, color, or development of rancidity. In the crude oils, particularly those from seeds, there are considerable quantities of *lecithins*,

$$\begin{array}{c} C_{17}H_{33}COOCH_2\\ \\ C_{17}H_{31}COOCH\\ O^{+} \\ |\\ (CH_3)_3 \begin{array}{c} NCH_2CH_2OPOCH_2\\ \\ - O^{-} \end{array} \end{array}$$

A Lecithin

cephalins, and other lipids which have not been completely identified.

$$\begin{array}{c|c} C_{17}H_{33}COOCH_2\\ \hline C_{15}H_{31}COOCH\\ \hline O\\ H_3NCH_2CH_2OPOCH_2\\ +\\ \hline O^-\\ \end{array}$$

A Cephalin

In freshly extracted corn or soybean oil, the phosphatides present may be as high as 2 or 3 per cent, but on purification most of these compounds

are removed. Sterols are

$$\begin{array}{c} \text{CH}_3 \\ \text{CH}_3 \\ \text{CH}_2 \\ \text{CH}_2 \\ \text{CH}_2 \\ \text{CH}_2 \\ \text{CH}_3 \\ \text{CH}_3 \\ \text{CH}_3 \\ \text{CH}_3 \end{array}$$

Cholesterol, A Common Animal Sterol

$$\begin{array}{c|c} CH_3 & CH_3 \\ CH & CH_2 \\ CH_2 & CH_2 \\ CH & C_2H_5 \\ CH_3 & CH_3 \end{array}$$

Sitosterol, A Common Plant Sterol

ilso present in crude oils to an appreciable extent. They vary from halibut iver oil with 7.6 per cent sterol to beef tallow with 0.08 per cent, with ourteen oils in the range from 0.1 to 1 per cent. The sterols occur in crude ats both as free sterols and as esters of fatty acids. They are partially renoved during the alkaline refining of the oil, but even in refined oil there is till some sterol. The phytosterol content of common vegetable oils is given a Table 2.2. "Phytosterol" includes all plant sterols with sitosterol the nost common. They constitute the major group of compounds in the non-aponifiable fraction of fats and oils.

Hydrocarbons are also present in some natural fats and oils. Squalene, is a highly unsaturated hydrocarbon with the formula

$$|(CH_1)_2C = CH(CH_2)_2C(CH_3) = CH(CH_2)_2C(CH_3) = CHCH_2 - |_2$$

nd occurs not only in fish liver oils but also in olive, rice bran, and nany others. Gadusene, $C_{18}H_{32}$, is named for the cod, Gadus, and was first olated from that oil. Pristane, $C_{18}H_{28}$, zamene, $C_{19}H_{38}$, and cetorhinene

are also in fish oil. The formula for zamene has recently been determined b Christensen and Sorensen⁴ to be

$$CH_2 = C(CH_3)(CH_2)_3 CH(CH_3)(CH_2)_3 CH(CH_3)(CH_2)_3 CH(CH_3)_2$$

2, 6, 10, 14 pentadec-l-ene

Waxes and some free fatty alcohols occur in very small amounts in fat and oils. The wax in corn oil has been reported to be composed of a mix

TABLE 2.2. PHYTOSTEROLS IN VEGETABLE OILS*

	· mg/100 g
Castor Bean	500
Cocoa Butter	170
Cocoa Butter	200
Coconut, Hydrog.	79.8
Coconut	60
Coconut	80
Corn .	580
Corn	1000
Corn Germs	800-1000
Cottonseed	311.2
Cottonseed	260
Crisco	232
Olive	133.7
Olive .	210
Peanut	247.9
Peanut	200
Peanut	190
Sesame	594.4
Sesame	520
Soybean	90
Soybean	380
Soybean .	195
Soybean	230
Wheat Germ	3600-6700

^{*}From Lange, W., J. Am. Oil Chemist's Soc., 27, 414-422 (1950).

ture of myricyl isobehenate and lignocerate, while another sample was reported to contain the esters of cetyl alcohol rather than myricyl.

C₂₁H₄₃COOC₁₄H₂₉ C₂₃H₄₇COOC₃₀H₆₁ C₂₃H₄₇COOC₁₆H₃₃
Myricyl Isobehenate Myricyl Lignocerate Cetyl Lignocerate

The coloring matter of fats and oils is composed of numerous pigments.

n orange-red or yellow pigmentation is usually caused by the presence of rotenoids which are soluble in oils. Carotenoids are highly unsaturated arocarbons, and when oils are hydrogenated, hydrogenation of the pigents also occurs with a reduction in color. Carotenoids are unstable at the temperatures, and if a fat or oil is treated with steam there is some as of color. They cannot be removed by oxidation since it brings the fat oil at or near to rancidity. They are, however, readily adsorbed on some sorbants such as fuller's earth.

The darkening which some oils show on limited oxidation is caused by e oxidation of tocopherols, the vitamin E substances present in some peopherol content of some fats is shown in Table 2.3. Occasionally when e oil is made from a green plant product, some of the chlorophyll will be tried along and color the oil. This is undesirable and very difficult to reove. Brown pigments are usually only present in oils which are prepared om rotten or damaged plant products and are believed to arise from the sintegration of protein and carbohydrate molecules.

We see, therefore, that most foods and food products contain not only be fats, but a great many related compounds. The classification "crude or even "fat" in food composition tables usually refers to the sum of ese compounds. Even purified fats and oils contain small amounts of mpounds other than simple fats, the triglycerides of fatty acids. The her compounds present are (1) the complex lipids—lecithins, cephalins, her phosphatides, and glycolipids; (2) sterols, both free and combined the fatty acids as waxes; (3) free fatty acids; (4) waxes; (5) pigments nich are lipid soluble; and (6) hydrocarbons.

IBLE FATS AND OILS

Much of our discussion will be confined to the prepared edible fats and swhich are sold in a fairly pure state. A great body of research has in built up around these foods because of efforts to differentiate one in the other so that a cheap oil is not sold for a high-priced one and because the manufacturers of shortenings have engaged in considere investigation. For many generations lard was the animal fat of choice preparing doughs and batters since it has sufficient plasticity at room because that it will cream with sugar and mix with egg or egg yolk, today in the United States the use of lard is small compared to the of "shortenings." Sometimes the shortenings are hydrogenated vegele oils: sometimes they are purified and standardized animal fats. The rket is highly competitive and during the years of development of se products much has been learned about fats.

Fatty Acids

Natural fats are mixtures of mixed glycerides in which the three fatt acids esterifying glycerol differ from each other. Little or none of th simple glycerides are present. Since the solubilities of these mixed glyceride are very similar, it is extremely difficult to fractionate them and to suc ceed in describing them in terms of the molecules present. After hydrolysi it is possible to separate the fatty acids; the available analyses of natura

2.3 TOCOPHEROL CONTENT OF FATS*

TABLE 2.3. TOCOPHEROL CONTENT OF THE	mg/100 g
	12.5
Cocoa	2.8
Cocoa	5
Coconut	8.3
Coconut	119, 102
Corn, Mazola	119, 102
Crude	104
Refined	83-110
Cot*onseed, Refined	110
Crude	3-30
Olive	26-51
Peanut, Refined	40-52
Crude	40-32
Safflower, Crude	18-65
Sesame, Refined	99-17:
Soybean, Refined	140-520
Wheat Germ	140-32
Influence of Processing on Cottonseed Oil	
Crude	102.8
Water Washed	102.0
Alk. Refined, Bleached, Filtered	100.
Refined, Bleached, Filtered, Deodor.	95.
Refined, Bleached, Filtered, Hydrog.	98.
Refined, Bleached, Filtered, Hydrog., Deodor.	97.

^{*}From Lange, W., J. Am. Oil Chemist's Soc., 27, 414-422 (1950).

fats are usually based on an analysis of the fatty acids rather than the actua mixed glycerides which occur in the natural product. Percentages of fatt acids by weight in some common fats are shown in Table 2.4.

Sometimes groups of fatty acids are separated. Molecular weight an the presence or absence of unsaturation affects (1) solubility in water, (2 volatility with steam, and (3) solubility of salts in water and alcohol. Thu fatty acid above lauric acid, $C_{11}H_{23}COOH$, is soluble in water even at 0 C. Butyric (C_4), caproic (C_6), caprylic (C_8), and capric (C_{10}) are volativity steam, while lauric (C_{12}) and myristic (C_{14}) are slightly volatile, he lead salts of the low molecular weight acids and unsaturated acids are ore soluble in ethyl alcohol than are the high molecular weight, saturated ids.

TABLE 2.4. PERCENTAGE OF FATTY ACIDS BY WEIGHT IN SOME EDIBLE FATS AND OILS*

tty Acids	Corn, %	Cotton Seed, %	Olive, %	Leaf Lard, %	Beef Tallow, %	Mutton Tallow, %	Butter, %
yristic							
C ₁₃ H ₂₇ COOH Imitic		1	1	1	2	2	10
C ₁₅ H ₃₁ COOH	6	21	9	28	32	34	30
C ₁₇ H ₃₅ COOH rachidic	2	2	1	8	15	19	11
C ₁₉ H ₃₉ COOH eic	1	1	1	_			—
C ₁₇ H ₃₃ COOH	37	25	80	56	49	43	30
C ₁₇ H ₃₁ COOH	54	50	8	5	2	. 2	3
C ₃ H ₇ COOH	_	_			adas (1971)		3
C,H,COOH	_	_			_	_	2
tal	100	100	100	98	100	100	99

*Adapted from Bailey, A. E., "The Chemistry and Technology of Food and Food Products," edited Jacobs, M. B., 1, 581–582, Interscience Publishers, Inc., New York, N. Y., 1944.

The groups of fatty acids are further separated into individual comjunds by esterification to form either the methyl or ethyl esters followed fractionation. They have been fractionated by very careful distillation at tremely low pressures, by crystallization from solvents at low temperares, by counter-current distribution in which they are partitioned belen two solvents, and by gas chromatography. Adsorption spectra with the separation of wave lengths is used to identify and even determine antities of fatty acids.

Gas chromatography is a technique in which the methyl or ethyl esters the fatty acids are passed over a column composed of a solid wet with methyl in which these esters will dissolve. A gas, usually helium,

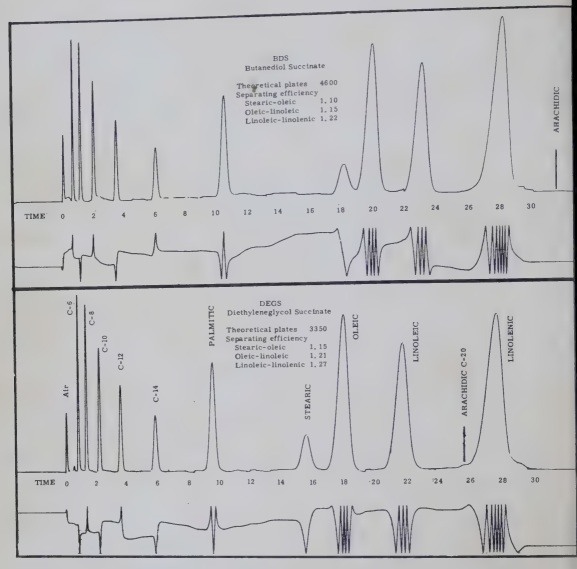


FIGURE 2.1. SEPARATION OF METHYL ESTERS OF FATTY ACIDS OF LINSEED OIL BY GAS CHROMATOGRAPHY.

Courtesy of Wilkens Instrument and Research, Inc.

then flows over the column and removes fractions. The most volatile will come off the column first.

The infra-red spectra of fatty acids and their esters have now been well established, and study of the spectrum of a fat or oil gives much information about the composition. High resolution of the lines in the infra-red portion of the spectrum is necessary in order that the presence of double bonds and cis or trans isomerism may be demonstrated.²²

The total unsaturated fatty acids are determined by methanolysis with methyl alcohol-sodium hydroxide followed by oxidation with standard potassium permanganate. The method has now been extended to include

re saturated fatty acids. After oxidation the acid oxidation products re removed by washing with alkali. The saturated methyl esters are recoved, dried, and weighed. Esters of fatty acids containing 16 or more arbon atoms with only small amounts of C_{14} are included among these sturated methyl esters. ¹⁸

$$R_{sat}.COOCH_{2}$$

$$RCOOCH + 3CH_{3}OH \longrightarrow$$

$$ICH=CH(CH_{2})_{n}COOCH_{2}$$

$$HOCH_{2}$$

$$R_{sat}.COOCH_{3} + HOCH_{2}$$

$$+ RCOOCH_{3} + HOCH_{2}$$

$$+ R_{1}CH=CH(CH_{2})_{n}COOCH_{3} \xrightarrow{KMnO_{4}} R_{1}COOH_{1}$$

$$+ R_{1}COOH_{2}$$

The fatty acids which are commonly present in natural fats are rericted to a surprisingly small number of possible compounds. The comon fatty acids are given in Table 2.5. Most of the acids are straightrain acids and, except for the low molecular weight acids, fatty acids are clusively composed of acids with an even number of carbon atoms. The isaturated acids have the possibility of either cis or trans isomerism, but nature only the cis isomer occurs. However, since formation of the ans isomers occurs when a fat is heated to a high temperature, is hydronated, or comes in contact with a number of catalysts, many fats and Is in commerce do contain trans isomers. In the unsaturated acids, 18 is a mmon number of carbon atoms and scarcely any fat lacks oleic acid. A mmon position for the double bond is between the ninth and tenth carons as in oleic, linoleic, and linolenic acids. When there are two or more ruble bonds, they are separated by a CH, group. Conjugation does occur a few fatty acids, but this is rare and occurs primarily in those fats ich are used as drying oils.

entification of Natural Fats and Oils

We will consider in this section the role of physical and chemical proper-

TABLE 2.5. MELTING POINTS OF COMMON FATTY ACIDS (MOST STABLE FORM)

No. of Carb Atoms			Meltin Point in
	Saturated		
4	Butyric	C ₃ H ₇ COOH	-7.
6	Caproic	C ₃ H ₁₁ COOH	-3.
8	Caprylic	$C_7H_{15}COOH$	16.
10	Capric	C ₉ H ₁₉ COOH	31.6
12	Lauric	$C_{11}H_{23}COOH$	44.2
14	Myristic	$C_{13}H_{27}^{\cdot}COOH$	54.4
16	Palmitic	C ₁₅ H ₃₄ COOH	62.9
18	Stearic	C ₁₇ H ₃₅ COOH	69.6
20	Arachidic	$C_{19}H_{39}COOH$	75.3
22	Behenic	$C_{21}H_{43}COOH$	79.9
24	Lignoceric	$C_{23}H_{47}COOH$	84.1
	Unsaturated		
16	Palmitoleic 9-hexadece		0.5
18	Oleic cis-9-octad	CH ₃ (CH ₂) ₇ CH=CH(CH ₂) ₇ COOH decenoic	16.3
18	Elaidic trans-9-oct	$CH_3(CH_2)_7CH = CH(CH_2)_7COOH$	43.7
18	Linoleic cis-cis-9-12	CH ₃ (CH ₂) ₄ CH=CHCH ₂ CH=CH(CH ₂) ₇ COOH 2 octadecadienoic	-5.0
18	Linolenic cis-cis-cis-9	CH ₁ CH ₂ CH=CHCH ₂ CH=CH(CH ₂) ₇ COO 9-12-15 octadecatrienoic	H -11.0
20	Arachidonic cis-cis-cis-c	CH ₃ (CH ₂) ₄ CH=(CHCH ₂ CH) ₃ =CH(CH ₂) ₃ COOH cis-5-8-11-14 eicosatetraenoic	-49.5
22	Erucic cis-13 cocos	CH ₃ (CH ₂) ₇ CH=CH(CH ₂) ₁₁ COOH	33.7

Physical Properties. The physical properties of the natural fats and oils e often used to identify them. Usually more than one property is easured so that the identification can be made with some assurance since dural fats and oils vary somewhat in their properties. Their composition not constant but varies slightly with climate, soil, and variety for vegeble oils, and with nutrition, season, and breed for animal oils.

Melting Point. Fats do not melt sharply but soften over a range of temcratures, and it is therefore impossible to apply the melting point techque, used in the identification of pure organic compounds, to them.

If a fat, a fatty acid, or some esters of fatty acids are heated very owly, they will melt, exist as a liquid as the temperature rises, and then blidify again. A second melting will then occur at a higher temperature. If we material is then chilled rapidly, it will melt at a lower temperature hen it is warmed again. This behavior has been known for many years, at it is only in recent times with the use of modern techniques that the colanation has been possible. Polymorphism, the occurrence of more than the crystalline form, explains this phenomenon.

Polymorphism is found in many long-chain carbon compounds. The umber of crystalline forms possible for each fatty acid or each ester is ill a matter of debate since many of the forms show melting points so ose together that it is difficult to separate them. However, for some combunds several crystalline forms have been established. These are designed alpha, beta, and gamma; the melting points of triglycerides are given Table 2.6. In compounds with several crystalline forms, one form is the lost stable and tends to be established.

Polymorphism is important to the understanding of the melting-point chavior of fats, fatty acids, and their esters. Furthermore, polymorphism plays a significant role in any operation where fats are solidized. In this latter instance environmental conditions must be controlled that the product will be uniform for example, when chocolates are apped, the formation of the high-melting forms is produced by controling the temperature of solidification.

When melting point is used to identify a fat, all aspects of the produre must be carefully controlled. A number of procedures have been de-

eloped and a few will be described briefly.

Solid fats are plastic over a fairly wide range of temperatures. By plastic e-mean that they are soft and can be deformed, but do not have the ability flow. The spreading quality of butter is the result of its plastic nature. her solid fats are examined microscopically, we see that they are composed of a mass of tiny crystals in a matrix of liquid fat. The crystals are of commeshed but are able to slide by one another and consequently give

TABLE	2.6.	MELTING	POINTS	OF	DIFFE	RENT	FORMS	OF
		SIMPLE	TRIGLYCE	RIDE	S IN	°C*		

Triglyceride	γ-form	α-form	β-form
Tricaprin	-15	18.0	31.5
Trilaurin Trilaurin	15	35.0	46.4
Trimyristin	33	46.5	57.0
Tripalmitin	. 45	56.0	65.5
Tristearin	54.5	65.0	71.5
Triolein	-32	-12	5.0
Trielaidin	15.5	37.0	41.5
Trierucin	6	25	32.5
Trilinolein	_	-43	-13.1

^{*}From Bailey, A. E., "Industrial Oil and Fat Products," 60, Interscience Publishers, Inc., New York, N. Y., 1945.

the mixed fat its plastic nature. A fat composed of only one kind of molecules does not possess this property of plasticity since it is, at any given temperature, composed either entirely of crystals or is liquid.

As a fat is warmed, the number of crystals distributed through the liquid fat diminishes and the amount of liquid increases so that the fat softens. If the number of crystals exceeds a critical amount, the fat will be hard and brittle and will lose plasticity. On the other hand, if the amount of liquid exceeds a critical level, the fat will flow.

Natural fats are complex mixtures of glycerides each with its own characteristic melting point. As the temperature of the fat is raised, the melting point of first one and then another of these glycerides is exceeded. Eventually a temperature is reached at which all of the glycerides have melted and the fat is liquid. The temperature at which this occurs is not sharply defined. The problem is further complicated by the fact that glycerides are soluble in one another. A particular type of molecule may dissolve in the liquid portion of the fat at a temperature considerably below its melting point.

Glycerides also have a tendency to supercool, i.e., to remain as liquids at a temperature below their melting point as they are cooled down from a higher temperature. The consistency of a fat at say 30°C may be different if it is heated slowly from room temperature than if it has been quickly cooled from a higher temperature. After standing, it will again come to equilibrium and the effects of supercooling will disappear.

The Softening Point of a fat is sometimes determined as a means of identification, but it cannot be applied to all fats. Capillary tubes are filled with oil and packed in ice over night so that the oil can solidify and come to equilibrium. The capillary tubes are clamped to a thermome-

r and submerged in a beaker of water. The temperature is slowly raised and the temperature at which the column of fat rises in the capillary tube called the softening point. The method gives reproducible results with time fats, rather poor results with others, while on lard compounds, for ample, it cannot be used.

The Slipping Point is another empirical method used to identify some atural fats and fat compounds. Small brass cylinders, filled with the solid it, are suspended in a bath close to the thermometer. As the bath is irred, the temperature is slowly raised. The point at which the fat rises the cylinder, or slips, is recorded as the slip point. The slip point is flated to the air or water beaten into the fat during its manufacture as ell as the composition of the fat. The slip point cannot, therefore, be recated on a particular sample with reproducible results.

The Shot Melting Point is the temperature at which a small lead shot will all through a sample. This method has some usefulness.

Why natural fats and oils differ in their melting behavior has been the abject of a great deal of study. How variations in the composition of a it and oil influence the melting has been studied in many free fatty acids nd pure glycerides. Likewise, analysis of the fatty acids present in fats nd a comparison with the melting point gives some information. In genral fats which contain relatively large amounts of unsaturated fatty acids ave relatively low melting points and are usually oils at room tempera-.re, while those with relatively large amounts of saturated fatty acids ave higher melting points. When the melting points of pure, simple triycerides are determined, it is found that lengthening the carbon chain f the fatty acids increases the melting point. For example, the melting oint of trimyristin (alpha form) is 46.5°C, tripalmitin, 56.0°C, and triearin, 65.0°C. Thus fats which are mixtures of glycerides of long chain, iturated fatty acids have higher melting points than those which have ther numerous unsaturated fatty acids or short chain ones, or both (see able 2.6).

Specific Gravity. The specific gravity of oils and fats is determined by the sual methods. The temperature is carefully controlled since significant nanges in these compounds occur in short ranges of temperature. The specific gravity of a fat or oil is usually measured at 25°C, but it may be necestry to use temperatures of 40°C or even 60°C for high-melting fats, anations in the specific gravity from one oil or fat to another are not cat. In general, either unsaturation of the fatty acid chains or increase in the length of the fatty acid residues tends to increase the specific gravity.

Refractive Index. The index of refraction is the degree of deflection of a cam of light that occurs when it passes from one transparent medium to

another. The refractive indices of fats and oils are often measured both because they can be rapidly and accurately determined and because they are useful in identification of these substances and the testing of their purity. An Abbé Refractometer with temperature control is used and the measurement is usually at 25°C. With high melting fats 40°C or even 60°C can be used, but temperature must be controlled and noted. The index of refraction decreases as the temperature rises; however, it increases with increase in the length of the carbon chains and also with the number of double bonds present.

Smoke, Flash, and Fire Points. The smoke point is the temperature at which a fat or oil gives off a thin bluish smoke. It is measured by a standard method in an open dish specified by the American Society for Testing Materials so that the evolution of smoke can be readily seen and reproduced. The flash point is the temperature at which the mixtures of vapor with air will ignite; the fire point is the temperature at which the substance will sustain continued combustion.

For a given sample of oil or fat, the temperature is progressively higher for the smoke point, flash point, and fire point. Table 2.7 gives data for a few oils with flash points measured in an open cup. The temperatures vary with the amount of free fatty acids present in an oil or fat, decreasing with increased free fatty acids. Since the amount of free fatty acids changes with variations in refining, the history of the oil or fat is important. The smoke point of a fat used for deep fat frying decreases with use of the fat. Fats and oils with low molecular weight fatty acids have low smoke, flash, and fire points. The number of double bonds present has little effect on the temperature required. Smoke, flash, and fire points are particularly useful in connection with fats used for any kind of frying.

Turbidity Point. The turbidity point of an oil is determined by cooling a mixture of it and a solvent in which it has a limited solubility. The mixture is warmed until complete solution occurs and then slowly cooled until the oil begins to separate and turbidity occurs. The temperature at which turbidity first is detectable is known as the turbidity point? The first solvent employed was glacial acetic acid in the Valenta test; but since it is difficult to keep the acid pure and since moisture has a marked effect on the test, other solvents have been substituted. In the Crismer test the solvent is methyl alcohol while in the Fryer and Weston modification of the Crismer test it is an equal mixture of 92 per cent ethyl alcohol and amyl alcohol. Data for some oils are given in Table 2.8.

The turbidity point determined for any one oil does show a range of values. It is particularly sensitive to the presence of free fatty acids and a

TABLE 2.7. SMOKE, FLASH, AND FIRE POINTS OF OILS*

					OILS	
	Smoke Points		Flash Points (Open Cup)		Fire Points	
Oil	°F	°C	°F	°C	°F	(
stor, refined	392	200	568	298	635	335
stor, dehydrated	348	176	570	299	638	337
rn, crude	352	178	562	294	655	346
rn, refined	440	227	618	326	678	359
iseed, raw	325	163	540	287	667	353
iseed, refined	320	160	588	309	680	360
ve, virgin	391	199	610	321	682	361
ybean, expeller, crude	357	181	564	296	664	351
ybean, extracted, crude	410	- 210	603	317	670	354
ybean, refined	492	256	618	326	673	356
-illa, raw	321	161	575	302	678	359
rilla, refined	352	178	608	320	685	363
rilla, refined	408	209	615	324	685	363

From Detwiler, S. B. and Markley, K. S., "Smoke, Flash, and Fire Points of Soybean and Other 2c able Oils," Oil and Soap, 17, 39 (1940).

rrection factor must be introduced for these acids. Nevertheless, different is show a wide enough range of values so that the test has value in the afterentiation of some oils and in the detection of adulteration.

Chemical Properties. A number of chemical tests have been evolved dure the years of study of oils and fats which are based on the partial dermination of the chemical composition of the oil or fat. These tests serve of the identify the fat and to detect the presence of adulteration. All oils id fats show some range of values; therefore sometimes more than one it is necessary. A few of the most commonly used tests are given below. Reichert Meissl Number is a measure of the amount of water-soluble latile fatty acids; the Polenske Number measures the amount of volatile soluble fatty acids; the Saponification Number, the amount of potassium droxide required to saponify the fat; and the Iodine Number, the amount unsaturation present. These chemical tests, then, differentiate fats and s on the basis of the chemical composition of the various triglycerides esent in the mixture.

The Reichert Meissl Number is defined as the number of milliliters of alkali (such as potassium hydroxide) required to neutralize the vola-water-soluble fatty acids in a 5 g sample of fat. The volatile acids will those in the range of molecular weights from butyric (C₄) to myristic acid. The Reichert Meissl test determines the amount of butyric and proic acids which are readily soluble in water and the caprylic and capric d which are slightly soluble. The Polenske Number is the number of

TABLE 2.8. CRISMER TESTS FOR VARIOUS	OILS	(FRYER	AND	WESTON) *
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Fat or Oil	Acidity (As Oleic) Per Cent	Observed Crismer Value	Correction Factor for Acidity	Corrected Value
Perilla	5.5	49.0	2.05	60.3
Linseed	2.0	58.3	2.05	62.4
Tung	0.9	74.0	2.05	75.8
Soybean	1.2	65.0	2.05	67.0
Niger	2.2	57.5	2.05	60.0
Sunflower	2.2	59.5	2.05	64.0
Corn	2.8	62.5	2.03	68.2
Cottonseed	0.1	65.0	2.03	65.2
Sesame	4.0	60.5	2.03	68.1
Rape	0.6	82.3	1.61	83.3
Almond	0.9	68.2	2.07	70.1
Peanut	1.1	72.0	2.07	74.3
Olive	0.7	67.8	2.07	69.2
Olive	1.8	65.5	2.07	69.2
Olive	3.6	61.5	2.07	69.0
Cacao	2.9	71.0	1.72	76.0
Chinese Vegetable Tallow	6.9	54.0	1.72	65.9
Palm	0.1	68.0	1.72	68.2
Lard	0.9	70.8	2.13	72.7
Tallow	0.1	72.5	2.13	72.7
Butter fat	1.9	43.0	1.54	46.0
Coconut	0.0	34.0	2.01	34.0
Coconut	1.6	30.0	2.01	
Palm Kernel	0.0	40.0	2.01	33.2 40.0

^{*}From Jamieson, G. S., "Vegetable Fats and Oils," 2nd ed., 36, Reinhold Publishing Corp., New York, N. Y., 1943.

milliliters of 0.1N alkali necessary to neutralize the volatile, water-insoluble fatty acids which are present in a 5 g sample. These two determinations are readily run on the same sample of fat and the Kirschner Value, described below, also can be carried out on the same sample.

Fat
$$\xrightarrow{\text{NaOH}}$$
 $\begin{cases} \text{glycerol} \\ + \\ \text{sodium soaps} \end{cases}$ $\xrightarrow{\text{H}_2\text{SO}_4}$ $\begin{cases} \text{Na}_2\text{SO}_4 \\ + \\ \text{free fatty} \end{cases}$ volatile $\begin{cases} \text{soluble} - \text{C}_4 \text{ and C}_6 \end{cases}$ acids $\begin{cases} \text{C}_4 \text{ to C}_{14} \end{cases}$ insoluble $\begin{cases} \text{C}_4 \text{ to C}_{14} \end{cases}$

A 5 g sample of fat is introduced into a flask and treated either with alcoholic sodium hydroxide or sodium hydroxide in glycerol. If alcohol is used, it must be removed by evaporation before the fatty acids are neutralized. Saponification occurs rapidly under the conditions of the experiment;

e soaps are neutralized with sulfuric acid, and the mixture distilled. On ndensation of the distillate the insoluble fatty acids precipitate. The stillate is filtered, an aliquot of the filtrate is titrated with standard tassium or sodium hydroxide and the Reichert Meissl Number calcued. The condenser, receiver, and filter paper are washed with water, and a insoluble fatty acids are dissolved in succeeding portions of ethyl alhol. This solution is titrated with standard alkali and the Polenske Numr calculated.

All of these determinations are particularly valuable in differentiating tter from coconut oil and in detecting adulteration of butter or the subtution of fat mixtures with the physical constants of butter for it. Only a fall number of other oils contain appreciable amounts of volatile fatty ids. The Reichert Meissl Number is particularly valuable in detecting ulteration in butter. Although the Reichert Meissl Number varies for tter with season, nutrition, and time in the lactational cycle of the cow, is usually between 24 and 34, higher than other edible oils. The distillation procedure as set up in the Reichert Meissl and Polenske methods does t remove all of the volatile acids from the saponification mixture; but if a procedure is followed accurately and the size of the sample restricted to a reproducible results and valuable information can be obtained.

The chief volatile, water-soluble fatty acid in butter is butyric acid. This letermined by the *Kirschner Value* which measures the potential amount soluble silver salts in the Reichert Meissl distillate. Silver butyrate is uble in water while the silver salts of the other volatile, water-soluble ty acids are relatively insoluble. The neutralized distillate from the inhert Meissl determination is treated with silver sulfate and filtered. The rate is acidified with sulfuric acid and distilled. The distillate is carely collected and titrated with 0.1N alkali, either sodium, potassium, or rium hydroxide. Kirshner Value is calculated from the following equant:

Kirschner Value =
$$\frac{A \times 121(100 + B)}{10,000}$$

cre:

A = corrected Kirschner titration, and

B = milliliters of alkali to neutralize the 100 ml distillate from Reichert Meissl.

equation results from the fact that the quantity of distillate collected in the 5 g sample in the Reichert Meissl (first) distillation is 110 ml while rul is titrated, and likewise in the Kirschner (second) distillation. But-

ter gives a Kirschner Value of 19 to 26, coconut oil approximately 1.9, palm kernel around 1, and other fats and oils between 0.1 to 0.2. Butter is there fore remarkable for the large amount of butyric acid present in it, and if can be identified on this basis.

The Saponification Number (the Koettstorfer Number) is defined as the number of milligrams of potassium hydroxide required to saponify 1 g of fat or oil. When potassium hydroxide reacts with a triglyceride, three moles of potassium hydroxide react with one mole of fat.

$$R_1COOCH_2$$
 R_1COOK $HOCH_2$
 $R_2COOCH + 3 KOH \rightarrow R_2COOK + HOCH$
 R_3COOCH_2 R_3COOK $HOCH_2$

If the triglyceride contains low molecular weight fatty acids, the number of molecules present in a 1 g sample of the fat will be greater than if the fatty acids have long carbon chains and high molecular weights. The fat with the low molecular weight fatty acids will consequently have a high Saponification Number. We find that butter with its unusually high percentage of butyric acid has the highest Saponification Number.

The method depends on the saponification of a weighed sample of fat of about 5 g with an excess of standard alcoholic potassium hydroxide. The potassium hydroxide is carefully measured into the flask with a buret or pipet. The mixture is boiled under reflux condensation until saponification is complete, and the remaining potassium hydroxide is determined by back titration with 0.5N hydrochloric acid.

The Saponification Number can then be determined by subtracting the number of milliequivalents of hydrochloric acid used for back titration from the number of milliequivalents of alcoholic potassium hydroxide introduced, multiplying by 56.1 mg (milliequivalent weight of KOH) and dividing by the weight of the sample in grams.

Saponification Number =
$$\frac{[(ml \times N \text{ KOH}) - (ml \times N \text{ HCl})] \times 56.1}{\text{g sample}}$$

The Hehner Value measures the amount of fatty acids which are insoluble in water. Thus fats and oils which have high Reichert Meissl Numbers will

ve low Hehner Values. Since most fatty acids present in natural fats are t soluble in water, most fats have relatively high Hehner Values. The me can be determined either by weighing a sample of fat and saponifying with alcoholic potassium hydroxide or by using the solution from the ponification Number titration. The alcohol is evaporated, the soaps disved in hot water, and treated with concentrated hydrochloric acid. Free ty acids are formed; and when the mixture is cooled, the insoluble fatty ids form a cake on the top of the solution. This is filtered on a weighed er paper, dried, and weighed.

The <u>Iodine Number</u> is the number of grams of iodine or iodine comund absorbed by 100 g of fat. The double bonds present in the unsatued fatty acids react readily with iodine or certain iodine compounds form an addition compound even while the fatty acid is combined with cerol in the fat. The Iodine Number is therefore a measure of the tent of unsaturation of the fatty acids present in a fat. While oleic acid ntains one double bond in its 18 carbon chain, linolenic acid contains ree double bonds in its 18 carbon chain. Thus a molecule of fat contain-2 one oleic acid can absorb or react with only one third as much iodine r IBr or ICl) as a molecule of fat containing one linolenic acid residue, the fatty acid assortment present in natural fats is fairly characteristic of 2 fat. While there will be variation in each vegetable oil with climate, al, and variety and in each animal fat with nutrition and breed, the rilations are small compared to the variations between fats. The Iodine 1 mber is therefore of great value in identifying fats and oils.

The Iodine Number is determined by dissolving a weighed sample of fat 1 g to 0.5 g) in chloroform or carbon tetrachloride and adding an exsolvence. After standing in the dark for a controlled period of time, excess, unreacted iodine is measured by thiosulfate titration. In the anus method the standard iodine solution is made up in glacial acetic acid d contains not only iodine but iodine bromide which accelerates the reion. The Wijs method uses an iodine solution made up in glacial acetic d but containing iodine chloride as accelerator. The excess iodine reswith sodium thiosulfate according to the following equation:

$$2 \text{ Na}_2\text{S}_2\text{O}_3 + \text{I}_2 \rightarrow 2 \text{ NaI} + \text{Na}_2\text{S}_4\text{O}_6$$

end point is determined by the disappearance of the blue starch iodine or.

The <u>Acetyl Value</u> is a measure of the amount of hydroxy fatty acids presimal fat. Castor oil is rich in ricinoleic acid, C₁-H₂(OH)COOH, and electermination is of value for this and a few other oils. Most edible and fats contain only very small amounts or no hydroxy fatty acids.

By the use of the physical constants and the chemical methods it is possible to differentiate and identify natural oils and fats. Some of these data are given in Table 2.9. Some variation occurs in all of the determinations because of the slight variations in composition of these natural fats. When it is said that a fat is characteristic of the species of plant or animal from which it comes, the statement is not true to an absolute degree. Nevertheless, by means of a few tests, an oil or fat can be identified and the presence of adulteration can be detected.

FLAVOR CHANGES IN FATS AND OILS

When fats and oils are stored, they undergo flavor changes which marked edly influence their market value. Consequently, a great deal of research has been devoted to an attempt to discover why some fats undergo changes more rapidly than others, what the changes are, what causes the changes and how they can be controlled. The flavor of rancidity is well known to everyone; and the problem of rancidity, although by no means completely understood, has been the subject of extensive research. Some oils and fats develop off-flavors before the onset of rancidity. This change is called reversion, and it, too, is important because it makes the fat or oil undesirable for food products.

Rancidity

It has been known for many years that fats and oils slowly take up oxygen for a period of time before it is possible to detect the flavor of the products of rancidity. This period is called the induction period, and it is followed by a second period in which the uptake is much more rapid. Rapid oxidation often continues for an extended period of time after which the rate falls off. The length of each period is markedly affected by many factors for each fat, and the course of the oxidation can apparently take a number of paths. Temperature, moisture, the amount of air in contact with the fat, light, particularly that in the ultraviolet or near ultraviolet, as well as the presence or absence of antioxidants and prooxidants influence the reaction.

Long ago farm women learned to store their lard in crocks with as small a surface exposed to the air as possible, and in a cool place. Darkness and coolness are not always possible when fats are shipped and stored during the ordinary course of commerce. Vegetable fats, particularly those from seeds, show a marked resistance to the onset of rancidity. Some seeds, if they are not bruised or crushed, can be stored for years without any change in the fats. But in general animal fats deteriorate fairly rapidly. Many years ago it was shown that the resistance of vegetable fats to oxidation rested

Stary	Specific Creaming IN 18 C	Solidification Point, C	Acid Value	cation Value	lodine Value	Meissl
Almond	0.914 to 0.921	15 to 20	0.5 to 3.5	183.3 to 207.6	93 to 103.4	0.5
Beef Fallow	5.895	31 to 38	0.25	196 to 200	35.4 to 42.3	0.25
Black Wahnut	0.918 to 0.921	turbid 12	8.6 to 9.0	190.1 to 191.5	141 to 142.7	
Butter Fat	0.907 to 0.912 40	20 to 23	0.45 to 35.4	210 to 230	26 to 38	17.0 to 34.5
Castor	0.960 to 0.967	turbid -12	0.12 to 0.3	175 to 183	84	4.1
Chicken Fat	0.924	21 to 27	1.2	193 to 204.6	66 to 71.5	∞ <u>.</u>
Coconut	0.926	14 to 22	2.5 to 10	253.4 to 262	6.2 to 10 2	6.6 to 7.5
Cocoa (Cacao) Butter	0.964 to 0.974	21.5 to 23	1.1 to 1.9	192.8 to 195	32.8 to 41.7	0.3 to 1
Cod Liver	0.922 to 0.931		5.6	171 to 189.	137 to 166	0.2
Corn	0.921 to 0.928	-10 to -20	1.37 to 202	187 to 193	111 to 128	4.3
Cottonseed	0.917 to 0.918 25	+12 to 13	0.6 to 0.9	194 to 196	103 to 111.3	0.95
Lard Oil	0.934 to 0.938	27.1 to 29.9	0.5 to 0.8	195 to 203	47 to 66.5	0.5 to 0.8
Linseed	0.930 to 0.938	19 to 27	1 to 3.5	188 to 195	175 to 202	0.95
Mutton Tallow	0.937 to 0.953	32 to 41	1.7 to 14	195 to 196	48 to 61	1
Olive	0.914 to 0.918	turbid + 2. ppt 6	0.3 to 1.0	185 to 196	79 to 88	0.6 to 1.5
Palm	0.924 to 0.858100°	35 to 42	10	200 to 205	49.2 to 58.9	0.9 to 1.9
Peanut	0.917 to 0.926	~	0.8	186 to 194	88 to 98	0.4
Perilla	0.930 to 0.937			188 to 194	185 to 206	ı
Poppy Seed	0.924 to 0.926	-16 to -18	2.5	193 to 195	128 to 141	0.0
Pumpkin Seed	0.923 to 0.925	-15	-	188 to 193	121 to 130	4.45
Rape Seed	0.913 to 0.917	01	0.36 to 1.0	168 to 179	94 to 105	0.010.0
Sesame	0.91925.	-4 to -6	8.6	188 to 193	103 to 117	1.1 to 1.2
Sova, Sov or Soja	0.924 to 0.927	10 to 16	0.3 to 1.8	189 to 193.5	122 to 134	0.5 to 2.8
Bean						
Sunflower	0.924 to 0.926	-17	¥1.2	188 to 193	129 to 136	0.5
Walnut	0.925 to 0.927	-15 to -27	2.5	190.1 to 197	139 to 150	0.92
Whale	0.917 to 0.924	-2 to 0	6.1	160 to 202	90 to 146	14

* Adapted from Lange, N. A., "Handbook of Chemistry," 14th ed., 678, Handbook Publishers Inc., Sandusky, Ohio, 1944

on the presence of antioxidants which occur naturally in the tissues and which are present in the oil when it is pressed. Some natural fats contain prooxidants which accelerate the onset of rancidity although most of the prooxidants, particularly the metals and their compounds, are picked up during processing from the metallic equipment.

The uptake of oxygen and the onset of rancidity seems to be related to the unsaturation of the fat, although this has been exceedingly difficult to show by direct comparison of natural fats. Since natural fats vary to a great degree in the occurrence of antioxidants, it is not surprising that contradictory results have been obtained. Studies of the autooxidation of simple esters of fatty acids have been more revealing. In one study, the relative rates of autooxidation under standard conditions were followed by measur-

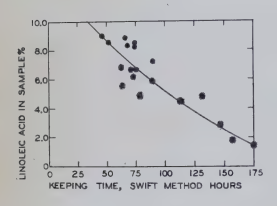


FIGURE 2.2. RELATIONSHIP BETWEEN LINOLEIC ACID CONTENT OF ALL-HYDRO-GENATED COTTON SEED OIL SHORTENING AND THE STABILITY OF THE SHORTENING. Reproduced from Bailey, A. E., "Industrial Oil and Fat Products," Interscience Publishers, Inc., New York, N. Y., 1945.

ing peroxide formation when oxygen was bubbled through the methyl ester: oleate, 1; linoleate, 12; linolenate, 25. Many other studies have been made, and it is generally agreed that the site of oxidation is the double bond.

The oxidation is not, however, a simple oxidation of the double bond. Many attempts to analyze the volatile compounds which can be smelled in a rancid fat show that a very complex mixture of compounds is formed. Heptyl aldehyde has been isolated most commonly and probably in largest amount. But heptyl aldehyde alone does not have the same quality of odor which is readily detected in a rancid fat. The compounds isolated have been relatively short chain compounds and include aldehydes, acids, hydroxy acids, ketones, and keto acids.

The products formed during the induction period have for many years been called peroxides and tests for the most part have centered around the ability of these compounds to release iodine from potassium iodide. This reaction with KI is very nonspecific, and whether or not the products are true peroxides has not been proved. In recent years hydroperoxides have been commonly named as the intermediates, and several theories have

een developed to explain their formation. A number of investigators have een able to isolate hydroperoxide from methyl oleate after oxidation at or ear room temperature.

9

The course of the oxidation is now believed to be a chain reaction and the current theory suggests that free radicals are the intermediates. It has been necessary to develop a theory which can explain how oleate with a couble bond between carbon 9 and 10 can yield products in which the oxyen is attached to carbon 8, 9, 10, or 11. It is suggested that the initiating eaction, under the influence of light, is the formation of a double free adical as oxygen adds to the double bond, one on the methylenic carbon and one on the oxygen. In the equation, unpaired electrons indicate the ite of the free radical.

A free radical may react with a methylenic carbon adjacent to a double cond to form another free radical:

$$RH + - \underset{8}{\overset{C}{\text{H}}} = \underset{10}{\overset{C}{\text{HCH}}}_{2} - \longrightarrow$$

$$RH + - \underset{8}{\overset{\dot{C}}{\text{HCH}}} = \underset{10}{\overset{C}{\text{HCH}}}_{2} - \text{ or } - \underset{8}{\overset{C}{\text{H}}}_{2} = \underset{10}{\overset{C}{\text{HCH}}} = \underset{11}{\overset{C}{\text{HCH}}} -$$

$$\uparrow \qquad \qquad \uparrow \qquad \qquad \uparrow$$

$$- \underset{8}{\overset{C}{\text{HCH}}} = \underset{10}{\overset{C}{\text{HCH}}} = \underset{11}{\overset{C}{\text{HCH}}} = \underset{11}{\overset{C}{\text{HCH}}} = \underset{11}{\overset{C}{\text{HCH}}} -$$

ecause each of the two free radicals formed is a resonance hybrid receiving contributions from two structures, there are four places at which an xygen molecule can add to the free radical:

$$R' + O_2 \rightarrow R-OO'$$

The free radical peroxide may then react with another methylenic carbor to form a hydroperoxide and another free radical:

$$R-OO' + R-H \rightarrow R-OO-H + R'$$

And so a chain reaction has started. With oleate four products—8, 9, 10 or 11 hydroperoxide form. Linoleate and linolenate are more reactive than oleate and this theory explains that reactivity. A methylene group between two double bonds will be more reactive and the possible number of isomers formed will be quite large.

$$-CH_2$$
 $-CH$ $=$ CH $+CH_2$ $+CH$ $-CH_2$ $-CH$

Each carbon in the portion of the chain shown is capable of forming a free radical and adding oxygen. Aldehydes and acids will form by cleavage of the carbon chain at the point of attachment of the hydroperoxide:

The course of oxidation in a fat is marked by a slow period of oxygen uptake called the induction period, followed by a much more rapid rate of absorption of oxygen. Figure 2.3 shows this change in rate of the uptake of oxygen by peanut oil as measured by the peroxide value. The graph also demonstrates the induction period, the time during which the antioxidant is undergoing oxidation. The sample of peanut oil used in this study was antioxidant-free. Graded amounts of antioxidant were added to subsequent samples. The series of curves show that as the amount of antioxidant increased, the length of the induction period increased. However, notice that the point at which the odor of rancidity products could be detected (arrows) was not constant for the various samples.

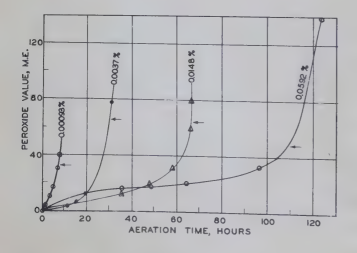


FIGURE 2.3. TYPICAL STABILITY CURVES OF FATS AERATED AT 99° C. Antioxidant-free fat containing different amounts of antioxidant concentrates (tocopherols) from peanut oil. Arrows indicate point at which oils smelled rancid. Reproduced from Bailey, A. E., "Industrial Oil and Fat Products," Interscience Publishers, Inc., New York, N. Y., 1945.

The presence in natural fats of minute amounts of compounds which rotect them from oxidation has been recognized for a long time. But the sentity of these compounds is still not entirely clear. The first compounds tentified were the tocopherols, the vitamin E's discovered by Evans and is coworkers. Four different tocopherols, α -, β -, γ -, and δ - tocopherol ccur. You will notice that α - tocopherol has three methyl groups in 5, 7, and 8 positions, while β - and γ - tocopherols have two (β - in the 5 and 8 ositions, γ - in the 7 and 8), and δ - only one. The tocopherols are readily xidized and consequently protect the fat from oxidation. The amounts of nese compounds present in refined oils are very small. (See Table 2.3, .18). Other antioxidants are believed to occur in natural fats since some ils with relatively low concentrations of the tocopherols are relatively table.

		Tocopherols	
	R_1	R_2	R_3
α	—СH ₃	—CH ₃	—СH ₃
β	—-CH ₃	—Н	—CH ₃
γ	—Н	—CH ₃	—CH ₃
δ	—Н	Н	—CH ₃

Numerous compounds have been developed synthetically and may be ided to fats to inhibit oxidation. Many of these compounds have been itented as antioxidants. The development of fats which do not require frigeration during the few months that they are stored in the ordinary tenen is chiefly the result of the use of these compounds. Some that have ten tried include the o- and p- dihydroxy benzenes, such as hydroquinone id pyrogallol; the aromatic amines; glucamine; gums; and cereal flours. The times antioxidants show a synergistic effect; two antioxidants may be an increase than would be expected from the sum of their activity. Butylated hydroxy anisole (BHA) or butylated hydroxy toluene (BHT) enften combined with propyl gallate. An example is a solution for proction of nuts, and composed of 14 per cent butylated hydroxy anisole, per cent propyl gallate, and 3 per cent citric acid in ethyl alcohol. A hall amount is used in nut candy.

The phosphatides, particularly the cephalins, seem to augment the antioxidant activity of other antioxidants such as the tocopherols. Since phosphoric acid and other acid products are capable of reinforcing the activity of some of the phenolic antioxidants, it may well be that the effect of the cephalins arises from the free hydrogen of the phosphate. Lecithins, however, do not have this effect.

The onset of rancidity in baked goods follows quite a different pattern from that of fat alone. The keeping quality of crackers and cookies is of great economic importance since these products are often stored for extended periods before they are consumed, and they are not protected from oxidation. Sugar in cookies and biscuits appears to have a marked inhibiting effect on the onset of rancidity. For example, McKinney and Bailey²⁰ found that biscuits made with hydrogenated vegetable oil became rancid in 80 days; whereas, when sugar was included in the recipe, they did not become rancid until 360 days. Oleo oil and hydrogenated lard gave similar results, although the difference with lard was not as marked. Rancidity occurred in the lard biscuits in 34 days, in those with added sugar in 50 days.

The effect of spices as antioxidants has been investigated, but the results are difficult to summarize briefly since considerable difference is obtained with differences in the food observed.³ For the most part spices, and especially cloves, are more effective in a water-fat emulsion than in dry lard and more effective in dry lard than pie crust. In a study of ground pork, mayonnaise, and French dressing it was shown that allspice, rosemary, cloves, sage, oregano, and thyme improved the stability of the fats. Clove has the greatest effect in ground pork, while oregano is most active in mayonnaise and French dressing. In oil-water emulsions, clove has the greatest activity, while in lard, sage and rosemary are outstanding.

Prooxidants promote the onset of rancidity. It is doubtful whether prooxidants occur naturally in fats and oils in significant amounts. Metallic ions act as catalysts in rancidity reactions, and they are often introduced in small amounts during the refining of the oil. If a small amount of rust is formed in steel equipment, it readily dissolves; or if copper vessels are used, small amounts of copper oxide may be dissolved. Since these metals are active at concentrations measured by parts per million, the nature of the utensils, pipes, and valves with which the oil comes in contact during refining is of great importance in its resistance to rancidity.

Tests for Rancidity. Rancidity tests have developed over the years in order to establish sensitive control of the stability of fats or the foods in which they are used. The more common tests are (1) Peroxide Value, (2) determination of carbonyl, (3) active oxygen determination, (4) thio-

parbituric acid test (TBA), and (5) Schaal oven test, which is used prinarily in the baking industry.

All of the tests for Peroxide Value measure the amount of iodine recased when potassium iodide reacts with rancid fat. The Lea method uses g of fat and 1 g of potassium iodide with an acetic-acid chloroform 2:1) solvent. After heating, the iodine formed is determined by titration with standard thiosulfate. Other methods modify quantities of solvent, but he principle remains the same.

The measurement of carbonyl compounds has followed traditional methods for this group. The <u>Kreis</u> and Shibsted tests were early ones which have dubious reliability. Today 2,4 dinitro phenyl hydrazine is commonly used in the Lappin-Clark method.

Active oxygen is a method which measures the length of time required o produce 20 meq peroxide per 1000 g fat when air is bubbled through ats under standard conditions. A bubble train is set up so that clean, try air bubbles through the oil at a constant rate such as 2.33 ml per sec. The oil is heated in a constant-temperature bath. Other samples are added at definite time intervals so that the time required for 20 meq can be calculated.

In the TBA method for determining rancidity, an oxidized or rancid fat sill react with 2-thiobarbituric acid (TBA) to form a red color the intensity of which is proportional to the amount of rancidity. In recent years this has been developed into a method for measuring the extent of rancidity in a ample of fat. In the rancid fat the compound formed which reacts with the 2-thiobarbituric acid is malonaldehyde, $CH_2(CHO)_2$.

The Oven Test is widely used in the baking industry. It takes little quipment and is very easy to set up. Biscuits, cookies, or crackers are tored in beakers or jars with loose fitting tops at 63°C or 145°F. The number of days required to develop rancidity is measured by odor and aste. The temperature is slightly above what might be encountered during istribution of the food through regular commercial channels. Its greatest se is in comparing fats. It is difficult to equate shelf life of a product with the Oven Test where conditions are variable.

teversion 5

Many oils and fats undergo a change in flavor before the onset of ranairty which is known as reversion. Some investigators believe that it is haracteristic of all fats, but that some fats show more pronounced hanges. The name "reversion" came to be applied to the change because one marine oils which possess a fishy flavor before processing revert to his fishy odor on storage. The term is poor because the flavors which develop in most fats were never there before, but it has become firmly established and cannot be displaced. The flavors which develop are quite different from rancid flavors. Soybean oil, which reverts readily, is described as developing first a buttery or beany flavor, then grassy or hay-like, then painty, and finally fishy.

The conditions under which reversion occurs are those encountered in marketing and also the high temperatures experienced during baking and frying. The problem of reversion is therefore of considerable importance in edible fats and oils. The factors which are known to influence the onset and development of reversion are (1) temperature, (2) light, (3) oxygen, to a limited extent, and (4) trace metals.

As the temperature of storage is increased, the length of time in which reversion flavors can be detected decreases. Thus Bickford found that soybean oil stored in the dark at 5°C (41°F) did not revert for several months, but oil stored at room temperature showed flavor changes in 10 to 14 days. The effect of light of various wave lengths on hydrogenated vegetable shortening was studied by Gudheim. He found that it is light in the blue-violet and from 325 to 460 m μ which causes reversion changes. Light at 325 m μ is in the ultraviolet and below this wave length does not pass through glass, so it is not a problem in ordinary packaging. Oxygen is not necessary in large quantities for the development of reversion, but a small amount appears indispensible since flavor changes which occur in oils stored in vacuum or under inert gases are not characteristic of reversion.

The effect of traces of metallic salts on reversion has been demonstrated by adding small amounts of salts to refined oils and measuring their tendency to revert. For example, soybean oil containing 0.015 ppm of iron was treated with iron salts to bring the levels to 0.03, 0.3, and 1.0 ppm. After storage at 60°C (141°F) for four days, the sample containing 1.0 ppm had the poorest flavor, 0.3 ppm intermediate flavor, and that with 0.03 ppm about the same flavor as the control. It has been shown that copper, cobalt, chromium, and zinc cause an acceleration in reversion. Aluminum, tin, and nickel are not so active. During the processing of fats, compounds are often added which have the ability to form complexes with metal ions. These are called "metal scavengers" since they remove the ions from the field of action and inhibit their catalytic effect. An example is EDTA (Ethylenediaminetetraacetic acid).

Rancidity and reversion are not the same thing. Indeed some oils and ts which are susceptible to rancidity, such as corn oil, are reversionistant. The flavor changes which occur during reversion vary with difrent fats, but in rancidity the final flavor is the same for all fats. The roxide value is widely used as a measure of the development of rancidity. it it is impossible to show a correlation between this value and reversion. The chemical reactions that occur in reversion are not completely nown, but much research has been done on the problem. It is still not vet finite exactly which compound undergoes change, what the change is, or hat products are formed. The flavorful products which impart the distinct wors of reversion are steam-distillable, and it has been demonstrated that ese products are present in reverted oils and fats in extremely small lantities. Numerous compounds have been suspected as reactants in reersion, but the data are impressive only for linolenic and isolinoleic acids. In 1936 Durkee, in searching for an explanation of the reversion tendnev of some oils and the reversion resistance of others, noticed that oils ith reversion tendency were particularly rich in linolenic acid. Dutton tered corn oil, which is naturally reversion-resistant, to a product with a gn linolenic content by interesterification with methyl linolenate in the resence of a catalyst. He also prepared a control by interesterification of e corn oil with methyl linoleate. After storage a taste panel compared ese two products with corn oil and soybean oil. The corn oil high in rolenate was judged as soybean oil (reversion-susceptible) in five out of x trials. This indicated that reactions leading to the flavor development in synthetically high-linolenate corn oil are similar to those in soybean oil. Some interesting work on linseed oil, which is not a food fat, has apiicated "isolinoleate" as the precursor of reversion. When linonic acid or its esters are hydrogenated, the first product of the adtion of one molecule of hydrogen is called "isolinoleic acid." vdrogenation of methyl linolenate produces 8, 14; 9, 15; and 10, 14 olinoleate.

$$C_{17}H_{29}COO + H_2 \rightarrow C_{17}H_{31}COO$$

Linolenic Isolinoleic

ns oils whose reversion tendency closely parallels the isolinoleate tent. As hydrogen is added and isolinoleate begins to form, the rersion tendency rises. This continues until the point is reached at ich hydrogen begins to add to the isolinoleate. As hydrogenation ntinues, the reversion tendency then falls. A concentrate of glyceryl-isolinoleate is very susceptible to reversion. In another study ethyl

linolenate was mixed with sunflower seed oil which is reversion-resistant and hydrogenated to shortening consistency. The sample reverted in pastry.

It appears as if both linolenate and isolinoleate may be responsible for reversion. The flavor of all reverted oils is not the same and even the same oil has different flavors under different conditions.

Some attempt has been made to identify the products of reversion which apparently result from degradation of linolenate and/or isolinoleate by steam distilling reverted oil and concentrating the distillate. Compounds which have been isolated and identified include 2-heptenal, $CH_3(CH_2)_3CH=CHCHO$, di-n-propyl ketone, $(CH_3CH_2-CH_2)_2CO$, maleic aldehyde, OCHCH=CHCHO, $\Delta^{2.4}$ -decadienal, $CH_3(CH_2)_4CH=CHCH=CHCHO$, and acetaldehyde, CH_3CHO . Exactly what contribution each compound makes to the subtle flavor blend in reversion has not been determined.

Much attention has been focused on practical methods of processing to prevent or minimize reversion. Avoidance of contamination with small traces of metals and the use of metal scavengers to minimize trace metal effects have been tried. Reversion is still an important problem in the refining and use of soybean oil, marine oils, rape seed oil, and a few others which have a marked tendency to revert.

THE TECHNOLOGY OF EDIBLE FATS AND OILS

Three principal methods are used for the extraction of edible fats and oils from the animal or vegetable tissues in which they occur. These are (1) rendering, which is chiefly applied to animal tissues; (2) pressing; and (3) solvent extraction. In general, in fats and oils there is more fatty animal tissue than vegetable tissue; in other words, less water, protein, and other nonfatty material are present. The problem in extracting the fat from any one tissue is always a little different than it is for another; and although a manufacturer of cotton seed oil may also make peanut oil in the same equipment, he is often limited to these two oils and does not attempt any other. Generalizations about methods of extraction of the fats can be made, but variations in procedure necessarily occur because of differences in the oil-bearing tissues. For details on methods of extraction of a particular oil or fat, see some of the books on fats.

Rendering

Rendering is a process by which fat is removed from a tissue by heat. It is also called "trying out." The tissue containing a high percentage of fat

carefully removed from the animal and chopped or minced. Heat allows e lipids to escape from the cells. If the heat is high, the cells are comtely ruptured, a cooked flavor develops, and "cracklings" are left with a oil floating on top. Rendering can be carried out either in the presence water—"wet rendering"—or in its absence—"dry rendering." In wet ndering the fat can be separated by gentle heat in an open kettle or in an toclave in the presence of steam. In the first method, the well-chopped sue is introduced into an open kettle along with a charge of water, rred gently, and heated to about 50°C. The fat floats to the top and is refully skimmed off. It has a bland flavor and requires little deodorization, but the process does not remove all of the fat from the tissue. It is uch more common today to use steam in digesters or autoclaves at relvely high temperatures and pressures of 40 to 60 psi. In this way the sue is quite well disintegrated and the separation of the fat is efficient.

Dry rendering is the process used in cooking bacon. The tissue is heated d the fat separates as the protein is denatured and the water is evapoted. Commercially the process is carried out under vacuum in steam-jack-

ed cookers.

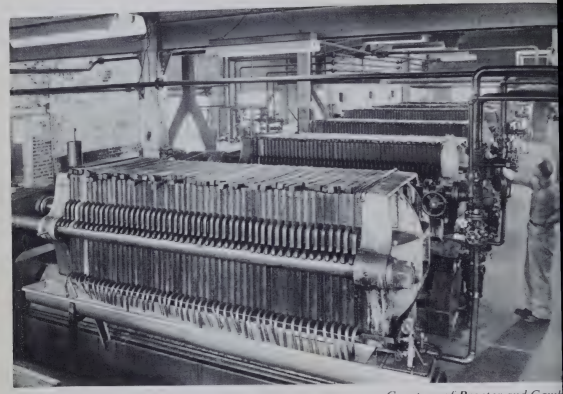
essing

Pressing is the application of high pressures to the oil-bearing tissue to uceze out the fat. In some cases, such as the pressing of olives, virgin is the first pressing of the fruit and is particularly bland in flavor. The fit is then subjected to subsequent pressings to give other grades of oil. Other oil products require that as much extraneous material as possible

removed from the seed or fruit before pressing. Thus cotton seeds are lintered and the kernel then separated from the hull, while in cereals the rm is separated before pressing is used. It is difficult to separate oils iciently by pressing and the greater the bulk of extraneous material, the wer the efficiency. Often the oil-bearing tissue is rolled, crushed, or bund to flaky particles. Usually the flake is cooked to denature the pron and release the oil. It is then pressed in a filter with filter cloths to ld back the extraneous material or it is passed through an expeller. An beller is constructed something like a meat grinder, with a worm screw an increases the pressure as the material is carried into it. At the end he worm is an opening for carrying out the residue with holes in the trom of the apparatus where the oil runs out. Sometimes the residue rm the expeller is then placed in a filter press.

Ivent Extraction

It's process has been used since Deiss extracted the press cake of olives



Courtesy of Procter and Gamb

FIGURE 2.4. PRODUCTION FILTER PRESS USED IN FAT AND OIL PROCESSING. In additional to simple filtration of raw materials, these filters are used to separate bleaching earth at the carbon used for decolorizing.

with carbon disulfide in Marseilles in 1855. In the United States solve extraction with petroleum ether has been used primarily for the production of soybean oil. (Garbage grease is also recovered by this method.) However, rapid improvements in continuous counter-current equipment has been such that in Europe solvent extraction is applied to many other oil In solvent extraction the tissue is treated in a fashion similar to the used in pressing to form a thin coherent flake. The oil is extracted eith in batch extractors in which the flake is gently agitated with successing portions of solvent, or by counter-current extraction where the flake are solvent move in continuous streams in opposite directions.

Common solvents are petroleum ether, benzene, chlorinated hydrocarbons, carbon disulfide, or Stoddard solvent. In the United States, petroleum ether boiling at 140°F to 185°F is used frequently, but hexarand other solvents are now coming into use. The solvent must then be moved from the oil. The method is quite efficient but is expensive because of the inevitable loss of the solvent through evaporation, and only in lar plants is it practical. However, in the removal of oil from tissues which have a relatively low percentage of oil, it is the only practical method

netimes a tissue is subjected to pressing and then the press cake with low fat content is extracted with a solvent.

horesidue from pressing or extracting is high in protein and is a value feed for cattle. If a solvent is used in extraction, this must be reved from the cake, of course, before it can be fed to cattle.

fining

he crude oils extracted from tissues often contain material that must be roved before these oils can be put on the consumer market or sent to the lrogenator. The crude oils may contain one or more of the following ups of substances: (1) cellular material or derivatives, both protein and bohydrate; (2) free fatty acids and phosphatides; (3) pigments; (4) odorcompounds such as aldehydes, ketones, hydrocarbons, and essential, and (5) glycerides with high melting points.

he first step in refining most oils and fats is the removal of finely died cell debris. This is usually accomplished by settling and then either tring or centrifuging. When the particles are colloidal, adsorbing or ering agents may be added before filtering.

ree fatty acids occur in some crude oils in goodly amounts. Thus palm usually has about 5 per cent free fatty acids. These can be fairly comely removed by steam refining and the remainder by alkali refining. nost all edible fats and oils are subjected to alkali refining since it re-



ourtesy of Procter and Gamble.)
URE 2.5. SOLVENT Ex10N. Mill room flaking

moves not only free fatty acids but also phosphatides and some otheresinous material.

Steam Refining. This consists of blowing steam through hot oil under vacuum. The free fatty acids with molecular weights below myristic steam distill but the fat is relatively nonvolatile. The process is used for the deodorization of fats. Oils with a high free fatty acid content are usually first subjected to steam refining; this is followed by alkali refining. Those with low fatty acid content may simply undergo alkali refining.

Alkali Refining. In alkali refining hot oil is treated with a solution of an alkali, usually sodium hydroxide (caustic soda) but sometimes with sodium carbonate (soda ash) and occasionally with other alkaline sodium salts. The free fatty acids react and form sodium soaps which are dispersible in the water layer. If the process is carried out rapidly, the amount of saponification of the oil that occurs will be very small.

In the United States, the common method of alkali refining is a continuous one in which the solution of alkali and oil is mixed, passed through a heater, and then through a battery of primary centrifuges. These centrifuges discharge the soap and alkali solution on one side and the of mixed with a small amount of soap and alkali solution on the other. The oil is then mixed with hot water and passed to a second battery of centrifuges that discharge washed oil relatively free of alkali or soap; the oil is then dried. Although alkali refining was once carried out by the batcomethod in which the alkali solution and oil are mixed in a conical kettle heated, and then separated, this process has been largely superseded by the continuous method which is more rapid and efficient.

Bleaching. Pigments are removed by adsorption on fuller's earth, other clays, and sometimes through the addition of small amounts of charcoal The process is called bleaching although it is a physical process. The oil is heated to 220°F to 240°F, agitated, and then filtered. It is returned to the kettle until the color of the filtered oil is sufficiently light. Some processor use vacuum since subjecting oils to high temperatures in the presence of air unfavorably affects their resistance to rancidity. Chemical agents, either oxidation or reduction agents, are not generally used for bleaching edible oils but only for industrial fats and waxes.

Steam Deodorization. The oils and fats demanded by the consumer toda must be very bland in flavor. By steam deodorization fats and oils can be rendered so bland that it is difficult to detect their origin by taste or smell Large kettles in which it is possible to maintain low pressures permit the use of only small quantities of steam for deodorization. The fat is introduced into the kettle and the pressure reduced to 1/4 inch or 6 mm of mer cury. The fat is heated and steam is injected into the bottom of the vesse. Oils are rapidly deodorized at 425°F to 475°F. If lower temperature



a. Soybeans.



b. Flaked beans ready for extraction.



c. Soybean oil.



d. Residual meal after grinding.

RE 2.6. STAGES IN SOLVENT EXTRACTION.

Courtesy of Procter and Gamble.

.sed, longer periods are required. The oil is then rapidly cooled. It is important to prevent the contact of hot oil with air, since oxygen is rious both to its keeping properties and to its flavor.

then cottonseed oil was first manufactured, it was noticed that the oil win from the tops of tanks in winter did not become cloudy or tend to liv, when refrigerated. Thus the term "winterized" came to be applied

to oils which have been processed so that they remain fluid at refrigerate temperatures. The process is applied to salad oils. The oils are slow chilled without agitation so that large, filterable crystals are formed. The crystals are composed of glycerides that contain the higher molecular weight and more saturated fatty acids and that consequently possess higher melting points. The cold viscous oils are carefully filtered and the filter calcused for some products in which the temperature of solidification is not in portant. Cottonseed oil with an iodine number of 107 yields 65 per cent can oil with an iodine number of 113 and a 35 per cent cake with an iodine number of 95. Lecithins can be added to oils to inhibit clouding an solidification. Thus a sample of cottonseed oil which clouds in 10 hours a 32°F will not cloud for 15 hours on the addition of 0.05 per cent soy bean lecithin.

Hydrogenation

Hydrogenation of fats and oils is the process by which molecular hydrogen is added to double bonds in the unsaturated fatty acids of the glycelides. Economically, hydrogenation is a most important process since by it use the physical properties of a natural fat can be altered. Thus the physical properties of the products can be regulated so that many natural fat can be interchanged, so that liquid fats can be substituted for plastic fats and so that an improvement occurs in the properties of natural plastic fats.

Hydrogenation occurs when hot oil saturated with hydrogen is brought in contact with an active catalyst, but the course of the reaction and it velocity is influenced by numerous factors. In most oils with a high content of C₁₈ unsaturated acids the possible reactions are (1) linolenic to linoleic or isolinoleic, (2) linoleic to oleic, and (3) oleic to stearic. When the ocontains unsaturated acids with carbon chains of other lengths, hydroge may be added to these chains. The products may be the natural linoleic anoleic acids or they may be the so-called iso acids in which the double bone is in a different position than that of the naturally occurring acid. Thus has been found that hydrogenation of linoleic may form natural oleic with the double bond in the 9, 10 position; but it will also form some iso-olei with the double bond in the 12, 13 position, which has a higher meltin point.

$$CH_{3}(CH_{2})_{4}CH = \overset{12}{CH_{2}CH} = \overset{10}{CH_{2}CH} = \overset{9}{CH_{1}(CH_{2})_{7}COO} - \overset{13}{CH_{3}(CH_{2})_{7}CH} = \overset{12}{CH_{1}(CH_{2})_{7}COO} - \overset{13}{CH_{2}(CH_{2})_{10}COO} - \overset{13}{CH_{3}(CH_{2})_{4}CH} = \overset{12}{CH_{1}(CH_{2})_{10}COO} - \overset{13}{CH_{2}(CH_{2})_{10}COO} - \overset{13}{CH$$

Methyl linolenate on hydrogenation gives 8, 14; 9, 15, as well as 10, 14 linoleate. Whale and fish oils contain rather large amounts of unsatuded C_{20} and C_{22} acids; and consequently, more variation in the molecules trogenated can occur. There is also the problem of the selective hydroation of fatty acid residues which occur on the α or β position of the zerol. Finally, during hydrogenation trans isomers are also often formed aer than natural cis isomers.

brouk and Brown¹⁸ examined the infra-red spectra of six commercial garine samples and five shortenings and found a high (22.7 to 41.7) per of trans isomers in all except one sample. O'Connor²² claims that rogenation of methyl oleate causes change to as much as 38 per cent of trans isomer. So the possible products are numerous and the complete rse of the reactions and the influence of all factors is not as yet known. 'evertheless, the course which the reaction follows is of great important as far as both the physical properties of the resultant fat and its sing qualities are concerned. It will be remembered that both linolenic isolinoleic acids have been implicated as active in promoting reversion tat the quantities of these acids remaining or formed influence the keep-

'hen hot oil and catalyst are stirred together under an atmosphere of ogen, the properties of the final product are affected by temperature, of mixing, nature of the catalyst, concentration of the catalyst, and sure of hydrogen.

quality.

number of active metals are capable of catalyzing this reaction, but strially nickel is used. Numerous methods are employed for producing incly divided metal with many active centers on its surface. Even when ethod of preparation is as carefully standardized as possible, to proactallyst with exactly the same activity in each batch is difficult. The catalyst is slowly poisoned during the hydrogenation, it needs to ractivated after it has been used for some time. Sulfur compounds, such drogen sulfide and sulfur dioxide which may contaminate the hydropoison the catalyst rapidly even when their concentration is low.



Courtesy of Procter and Gam

FIGURE 2.7. EXTERIOR OF EQUIPMENT USED IN HYDROGENATION OF FATS.

catalyst in a vacuum. It is also a contaminant of hydrogen made by the steam-iron method.

Hydrogen of reasonable purity is desirable to protect the catalyst. the United States the hydrogen used in this process is made by electrolys in areas where electric power is relatively cheap. This gas is quite pur In other areas the hydrogen is made by the steam-iron method. Iron ore alternately oxidized by steam with the formation of hydrogen and then reduced. The reducing agent is usually "blue gas" or "coal gas," a mixture hydrogen and carbon monoxide made from hot coke and steam.

$$C + H_2O \rightarrow CO + H_2$$

 $3Fe + 4H_2O \rightarrow Fe_3O_4 + 4H_2$

Hydrogen synthesized by this process is purified to remove sulfur con

nds formed from the coke, as well as carbon monoxide and carbon vide.

he hydrogenation is usually carried out by the batch method in large els with a capacity of 10,000 to 25,000 lbs of fat. The vessels are pped with heating and cooling coils. Oil and catalyst are combined and ed under vacuum between 150 F and 400 F. Hydrogen is introduced er pressure in such a manner that the maximum amount is dissolved in oil. Variations in the apparatus are not only designed to improve the ation of the oil but also to facilitate the introduction of hydrogen. The perature and pressure used vary with the oil being hydrogenated the characteristics of the final product. Sometimes the hydrogenation arried out in two stages at different temperatures. The reaction is ed when the fat has reached the desirable iodine number or consistency. line number and consistency parallel one another when the procedure is dardized; otherwise they do not.) The tank is evacuated and the oil ed to about 150°F. The hot oil must not come in contact with air, ourse. The oil is filtered to remove the catalyst and chilled rapidly so small crystals are formed. Sometimes monoglycerides, antioxidants, other compounds are added, depending on the product that is desired. is sometimes whipped into the product to impart a snow-white appear-

hnology of Individual Fat Products

everal groups of fat products are as follows: (1) butter, (2) oleo, oil, and stearin, (3) lard, (4) salad, cooking, and frying oils, (5) shortenings, and nargarine.

although some is still produced in farm kitchens. The tendency is to creamery depots that collect cream or milk from the farms and then is to the factory. Cream varies greatly in its quality and is usually the on a graded scale that is based on acidity, flavor, odor, and the unit of foreign material present. High-quality butter cannot be made putrid, dirty cream. A small amount of butter is churned from sweet n, but by far the greater part is made from ripened cream. Salt is lly added to the extent of 2.5 to 3.0 per cent. The minimum butter from its established by law at 80 per cent; and the remainder is posed of buttermilk, water, and milk solids.

e common practice is to neutralize the cream with sodium bicarte magnesium oxide, or calcium carbonate: pasteurize it to kill organisms; and inoculate it with a starter of selected bacteria. After in 1 for 3 or 4 hr, during which time the bacteria grow and produce

flavorful compounds and acids, the cream is churned. The butter separa and is washed with cold water. It is then kneaded to reduce the water co tent, salt is added and blended in, and the product is packaged.

The use of a starter for the ripening of the cream has made possible of production of a high-flavor, standard butter. It has been found that of flavor of butter is the result of a number of compounds, but the most is portant is diacetyl. This compound is synthesized by the action *Streptococcus citrovorus* and *Streptococcus paracitrovorus* on citric as present in the cream. It has been shown that acetyl methyl carbinol is an termediate in the synthesis and that both compounds are present in the ripened cream and butter.

 $CH_3COCH(OH)CH_3 \rightarrow CH_3COCOCH_3$ Acetyl Methyl Carbinol Diacetyl

Lactic acid and small amounts of propionic and acetic acids are form from fermentation of lactose present in the cream. Streptococcus lactis added to the starter along with Streptococcus citrovorus and Streptococcus paracitrovorus so that these reactions can be assured.

The consistency of butter varies somewhat during the year with variati in feed. High fat diets of oil cakes have a particularly marked effect on to composition of the fat secreted in the milk. The age and breed of the care also of some importance in determining the composition of the greenides present and consequently of the consistency of the butter.

Oleo Oil and Oleostearin. These substances are prepared by fraction tion of the fresh, selected, internal fat tissues of beef. The tissues a carefully rendered at low temperatures and the fat produced held at about 90°F for several days until partial crystallization has occurred. A sle crystallization is desirable so that large, easily filtered crystals will formed. The mixture of crystals and oil is then filtered in a filter press yield oleo oil and cakes of oleostearin. Oleo oil is used in the manufacture of some margarines and also unmodified as a shortening by confectioned and bakers. Oleostearin is used in the manufacture of compound shorterings by blending with other fats and in the production of some special margarines.

Lard. This substance is the processed fat of hogs. The quality of the la and its characteristics depend on the part of the carcass from which it taken, the way in which it is processed, and the feed of the animal. Le lard is the highest quality lard; it is rendered from the leaf fat (the fat tissues around the kidneys). Fatty tissue from the back of a hog gives so ond quality lard, while that from the intestines is lowest in quality. T

may be either dry or wet rendered. A great deal of the lard produced to United States is steam rendered.

reresterification in Lard. ... A recent development in the technology of may return this fat to its former position as a widely used and highed fat. This is directed interesterification of lard which improves its tic range and diminishes its tendency to graininess. Natural lard tends evelop fairly large crystals of disaturated glycerides (fats with two satu-I fatty acid residues) that give a grainy texture when the lard is chilled make it difficult to cream in batters and doughs. Since the melting ts of the disaturates are in the range of room temperature, the lard ens so much that it has little body and almost no plasticity at these peratures. Lard has approximately 37 per cent saturated fatty acids, cipally stearic and palmitic. Under suitable conditions the glycerides ent undergo interesterification with the rearrangement of the fatty acids e glycerides. In directed interesterification the reaction is carried out temperature where all the glycerides exist in the liquid phase except risaturated esters that are solids. As trisaturated glycerides are formed, crystallize and are removed from the reaction mixture. The interesteriion no longer is random, but is driven toward the formation of these trisaturates. The catalyst for the reaction in commercial interesterifiin is an alloy of potassium and sodium.

and is now interesterified commercially by a continuous process in a the catalyst is metered into the fat in very small particles. The temture is carefully controlled and the fat is constantly agitated as crystal-on occurs. The level of trisaturated glycerides formed can be conceed by the temperature of the fat and the length of time allowed for allization. As the percentage of trisaturates increases, the resistance of and to softening at room temperature is increased or, expressed in her way, the percentage of solids at a given temperature increases.

e catalyst is quenched by reaction with water and carbon dioxide. The int of hydrolysis of fat is minimized by the introduction of carbon de along with the water, which prevents the pH from rising too high. soap formed is removed by centrifugation, washing, and further if gation. The lard is then dried in a continuous vacuum drier.

crease its consistency. It is deodorized and supplemented with antiunts and monoglycerides before packaging.

hough the process is applied commercially to lard, interesterification natural fats will occur at elevated temperatures. Often the rate of on is slow and a considerable period of time is necessary to reach baum. The change in the composition of the glycerides that is

brought about by interesterification changes the physical properties of t fat, of course. After interesterification cocoa butter has a much high melting point and a very different consistency. Some fats are more radical altered than others. Many catalysts have been used but the alkali met and their alcoholates are most active.

Salad, Cooking, and Frying Oils. These are prepared oils and, with the exception of virgin olive oil, they are alkali refined, bleached, winterize if necessary, and deodorized. Salad oil is an oil which will remain sustantially liquid at a temperature of 40°F to 45°F and forms a mayon naise emulsion stable at these temperatures. An oil labeled "cooking of will not have these properties. Most oil is sold in the United States salad oil. The oils on the market in the United States are principally of tonseed oil, followed by corn, soybean, olive, peanut, and rather small quantities of sesame, sunflower seed, safflower and others. Cottonseed must be winterized before it can be sold as salad oil. Some corn oils a soybean oils deposit small amounts of crystals at refrigerator temperature if they are not winterized. Peanut oil forms very small crystals that a difficult to filter, and it is therefore not easy to winterize.

In the United States, where factory methods are used for the production of fats and oils, the extent of extraction of the crude oils is high and flavor is strong. Since the consumer market demands oils with bland flavor oils must be deodorized before they are packaged. Olive oil is the ception, since it is customary to anticipate a flavor in olive oil. The ol oil is usually blended so that the variations in season and climate that aff the flavor can be minimized and a reproducible flavor obtained. The mount of the United States is, of course, relatively small.

Bleaching is likewise necessary because of the demands of the consum Most salad oils on the market are relatively light in color. A darker of tonseed oil is produced for a section of the market that is accustomed the darker color of olive oil.

Oil that is packaged as cooking oil is sold for commercial deep fat fing. In hotels and restaurants almost all frying is of this kind. Also, in preparation of doughnuts, deep fat frying is used. The oils sold for to purpose are alkali refined and deodorized but are sometimes not bleach since a light color is not important. Potato chips and products not to consumed immediately are usually fried in solid fats rather than cook oils.

Mayonnaise is an emulsion of water in oil that readily breaks if cryst begin to form in the oil. Therefore, the oil used in mayonnaise and sa dressing should be thoroughly winterized. The oil is alkali refined a deodorized, but the amount of bleaching may be small. Mayonnaise is

ed to have a yellow color and the use of a dark oil in its preparation erefore not objectionable.

cortenings. Two types of shortening may be found on the market in the ed States: compound shortenings, which are mixtures of high-melting such as animal fats and low-melting fats, hydrogenated fats, or oils; those formed by the hydrogenation of an oil or a mixture of oils to the red point. Compound shortenings now comprise only a small bulk of shortenings sold in the United States. Compounding first began in the 1s when beef fats were added to lard. Later cottonseed oil was addedally, although the products were still sold as "lard" or "refined lard," amount of lard in them was sometimes small. After a Congressional integation, the use of the term "lard compound" was made mandatory, ntually the term "shortening" was introduced; since it has no implicate as to the origin of the fat but only as to its use, this term has befer firmly established.

This arose because the early settlers were from northern Europe where er and lard were the fats most commonly used. The cooking practices outhern Europeans, with their use of liquid fats such as olive oil, were oduced later when migration occurred in the nineteenth century. This not alter the basic pattern of American cookery but simply expanded enriched the cuisine, making it more cosmopolitan.

1900 all shortenings on the market were attempts to imitate lard as ally as possible. But with the introduction of hydrogenation about 1910, ufacturers of all-hydrogenated oils, usually cottonseed oil, began to graze the advantage of developing their ware as a new product and not ply as a substitute for lard. The shortenings on the market today are inferent from lard. They have much greater resistance to rancidity and petter creaming or emulsifying agents in batters. They are completely d in flavor; and although this may not be an advantage to the tongue of a concurr, the American consumer has come to expect all fats except and oleomargarine to be flavorless. As a result, the use of lard has cased and today lard is being processed in some plants to resemble the ogenated shortenings.

mimber of oils and fats that can be used for the production of chings is quite large, since a suitable consistency can be obtained by ang various hard and soft fats or by hydrogenating soft fats or oils. Solid or hard fats used in compounded shortenings are oleostearin or e allow as well as lard blended with various oils, principally cottonioil. When soybean, whale, or fish oils are used, they are usually parhydrogenated in order to reduce their marked tendency to flavor re-

version. Hydrogenated all-vegetable shortenings are made principally foottonseed, peanut, and soybean oil in the United States, although amount of soybean oil is limited by its tendency to reversion. Cotton oil is the principal oil, but does not yield a hydrogenated product wit long a keeping period as peanut oil.

Usually the oils are carefully blended and the hydrogenation control so that the glycerides will form a product with the longest possible kee period, plasticity over a long temperature range, and a consistency makes for ready incorporation into doughs and batters. Since resistant oxidation and the appearance of rancidity is related to the amount linoleic acid present in the final product, the amount of this acid, inso of oleic acid, should be reduced as much as possible during hydrogenated However, hydrogenation of linoleic acid yields not only oleic acid also iso-oleic acid, which has a tendency to harden the fat and reduced plastic range. A balance must be achieved, therefore, in which both line and iso-oleic are kept at a minimum. Some manufacturers produce shortening by blending two different hydrogenated fats; others control properties of the product during hydrogenation and do not use blending

Special hydrogenated shortenings are prepared for the bakers of subsecuits, cookies, and crackers. A long shelf life for the products me from these special shortenings is absolutely essential; the fat of all of ingredients is more likely to undergo flavor changes either from rever or rancidity. Most of these products are prepared in factories and me in large batches where the temperature can be controlled. It is not essent for shortenings used under these conditions to be plastic over a wide to perature range. Therefore, biscuit- and cracker-type shortenings have maximum resistance to oxidation and a short plastic range.

Since 1934 superglycerinated or "high ratio" (this term is copyrigh by Proctor and Gamble Co.) shortenings have been on the market. The are shortenings to which monoglycerides and diglycerides have been ad to promote the emulsification of fat in water and to increase the amount water and consequently the amount of sugar that can be incorporated batter and still produce a cake with sufficient strength not to fall durbaking or cooling. Formulas for each of these two types are shown below

monoglyceride or diglyceride is both hydrophilic, because of the hydroxyl groups, and lipophilic, because of the fatty acid residues. In r words, part of the molecule has an attraction for lipids but none for r; whereas part possesses an attraction for water. Consequently, these pounds are able to promote emulsification, to act as emulsifying agents t and water mixes. The practical result is that recipes can be used in hith amount of liquid is increased and the ratio of sugar to flour is. This produces a sweeter cake. When monoglycerides and diglycerides idded to a shortening it is possible to prepare cakes with a sugar to ratio of as much as 1.4:1, while with ordinary shortenings a ratio of annot be exceeded. This is the origin of the term "high ratio shorten-

onoglycerides and diglycerides are prepared by treating a fat with one to one sixth its weight of glycerol and adding a small amount of son hydroxide as catalyst. The reaction mixture is heated with stirring an atmosphere of an inert gas. When the reaction is complete, it is ed with an excess of phosphoric acid, dehydrated, and filtered to recthe fat-insoluble sodium phosphate. The product contains a mixture onoglycerides, diglycerides, and triglycerides, a small amount of free rol, and some free fatty acids formed by the neutralization of the with phosphoric acid. This mixture is added to the shortening at the of deodorization.

readdition of the monoglyceride and diglyceride mixture to shortenincreases their price by a small amount but also increases their useis in doughs and batters. They are therefore widely sold for this ose. When the shortening is used for deep fat frying, the addition of compounds is actually detrimental since they decompose readily and the smoke point. Antioxidants are commonly added to shortenings crease their resistance to rancidity.

nsumption during the past few years. It was first developed by the higher than the higher than

other fats. The addition of milk and the udder tissue probably resulted the introduction of microorganisms that soured the milk and production of those found in butter made from sour milk. The incorporation of the milk with the oleo oil also produced a product while like butter, is an emulsion of water in fat. The name "oleomargarity was based in the belief at that time that the fraction of beef fat used to straight chain, C_{17} acid, which has not been found to occur natural Through the years many changes have occurred in the manufacture margarine, but the basic process is still the same.

Today margarine is prepared from a variety of fats and oils. Oleo oil a lard—as well as the same vegetable oils that are suitable for high gr shortenings, i.e., cottonseed, peanut, sesame, palm oils, etc., and coco oil—can be prepared in bland form. The amount of soybean oil must limited because of its tendency to reversion, and fish oils are not suita for the same reason. In Europe whale oils are used for margarine ma facture. The fats must be carefully extracted and refined so that the firm product does not have a detectable flavor from the oils but only from milk incorporated in it.

The consistency of margarine is of great importance in the success of product. It must melt in the mouth as butter does, since a residue leave pasty sensation. Margarine is definitely a substitute for butter and, in eyes of the consumer, must have properties similar to it in order to acceptable. Margarine must be quite plastic at room temperature so that spreads readily and be fairly hard at 40°F to 45°F refrigerator temperatures, as butter is. In the temperature range from 45°F to 60°F butte too hard to spread easily, and much margarine is now produced which superior to butter in its plasticity in this range. The consistency of margarine is the result of the fats used in its preparation, the extent of hydrogenation, and the course of the reactions during hydrogenation. Usual margarines are produced by carefully controlling the hydrogenation of total body of fat rather than by blending.

A few specialty margarines are manufactured, and the consistency these products does not require the rigid specifications demanded by ta margarine. Margarine for puff-pastry, for example, has rather a high m ing point and is tough and waxy at room temperature, so that the roll in of the fat into the dough is facilitated.

In preparing margarine the natural fats and oils are carefully extract alkali refined, deodorized, and then hydrogenated to the desired consency. The fat is then emulsified with ripened milk. In the United Staskim milk is commonly used. It is pasteurized to destroy bacteria and the



Courtesy of Kraft Foods.

FIGURE 2.8. MARGARINE INGREDIENTS ARE AUTOMATICALLY MIXED IN THESE TANKS.

oculated with a strain of select bacteria that can produce compounds the desirable flavors in the milk and in the emulsion. These are the same ans of microorganisms that are used in the production of butter. The oculated milk is held for 12 to 24 hr to permit the growth of the organism. The ripened milk is run into the liquid fat and stirred vigorously. The ripened milk is run into the liquid fat and stirred vigorously.

Emulsifying agents stabilize the margarine and prevent leakage, the seption of fluid during storage. They also prevent the rapid separation of and water when the margarine is melted, spattering, and the sticking of lk solids to the bottom of the pan. In butter natural emulsifying agents present that hold the water in the emulsion and when the butter is ted, allow steam to escape by foaming rather than by spattering.

Lenthins, particularly those from soybeans, are widely used as emulsing agents in margarines. A number of synthetic products are also used, e mono and diglycerides used in the formation of superglycerinated prenings help stabilize the emulsion and prevent leaking, but do little to event spattering.

The sodium sulfoacetate derivative of mono and diglycerides are effective in minimizing spattering and are added to many margarines for this purpose. The fat-milk emulsion is cooled and the plastic, solid mass held for some time to allow bacterial action and the development of flavor. Salt is then added to the extent of 2.5 to 3 per cent of the total weight. Since the salt dissolves in the aqueous phase, the salt content of these tiny drops is much higher. It is so high that the activity and growth of the bacteria is stopped. The margarine is worked or kneaded during the operation of salting and the crystals are reduced so that no graininess occurs.

In the United States most margarine produced for the consumer marke is fortified with vitamin A or provitamin A, the carotenes, to the extent o 9000 units per pound. A yellow dye is added to much of the margarine sold in this country since it has become legal to do this without the pay ment of a high excise tax. Sodium benzoate is occasionally added as a preservative.

The Shortening Value of Different Fats

Baked goods all use fat as one of their ingredients and its role in the cookery process is particularly important in those leavened by baking pow ders or those of the pound-cake type that are leavened by air. The amoun of fat in most baked goods is considerable: in pie crust, 15 20 per cent pound cake, 15–30 per cent; doughnuts, 15 20 per cent; cookies, 5–20 pe cent; cake, 10 20 per cent; crackers, 5 12 per cent; but in bread only ap proximately 1 per cent. The fat is not only important for the flavor it im parts but also in the development of texture and the tenderness of the product.

In pound cake leavening occurs by incorporating air into the fat during the creaming process. When the batter is heated in the oven, the small ai bubbles expand and fill with steam. Indeed, the data shown in Table 2.10 indicate that the greater percentage of leavening in pound cake come from the steam that collects in the tiny air bubbles rather than from the air itself. The other components, principally the gluten in the flour, form the walls around each little bubble, and during baking they set to a fairly rigid structure. When the cake is removed from the oven and is cooled, the air in the small bubbles contracts and the steam condenses. But in a well balanced recipe the walls are sufficiently strong to hold up and the cake does not fall. The role of the fat in the leavening process is in its ability to trap air during the mixing process. The aqueous phase composed of the milk and eggs and dissolved substances such as sugar and salt does no have this ability. So the fat is indirectly responsible for the texture of the cake. Its tenderness is likewise believed to be a function of the fat. The

alls around each bubble are made of gluten and starch which can be very sugh and hard. Fat is streaked through and serves as a lubricant so that hen the cake is bitten, particles of gluten and starch slide on one another id the walls crumble.

In butter cakes fat has much the same role as in pound cakes although avening does not depend on air and steam alone but is supplemented by e carbon dioxide produced from baking powder. If the batter is formed, creaming the fat first, the air bubbles may serve as the nucleus for carmidoxide bubbles. Cakes with satisfactory grain can be produced, hower, by other methods of mixing. Certainly the fat in butter cakes is important as far as tenderness is concerned and allows the gluten particles to de on one another when the cake is bitten or cut.

In pastry a stiff dough is rolled out. The fat is flattened into sheets tween the layers of flour and water and causes their separation. This prouces flakiness. The fat is also squeezed in between gluten particles or
ands so that a continuous tough matrix is not formed but rather one
nich is very brittle or short. This also occurs in high-fat cookies.

Fats differ considerably in their ability to render a cake tender or a stry or cookie short. In commercial bakeries it is essential that products uniform. A marked variation in the fat may have disastrous effects on e bakery goods.

The relative shortening value of a fat is determined by means of a prometer. This apparatus measures the load necessary to break a wafer epared under standard technique and with a standard recipe. If it takes a

TABLE 2.10. TYPICAL EFFECT OF INCORPORATING AIR IN A POUND CAKE FORMULA CONTAINING 21 PER CENT FAT AND NO CHEMICAL LEAVENING AGENT*

В
w high
7 377
3 662
6 285
0 1520
7 858
2 71
7.0 8.3
3.0 91.7

Fr. Bicles, A. F., "Industrial Oil and Fat Products," 278, Interscience Publishers, Inc. & York, N. Y., 1945.

very small weight to break the wafer, then the wafer is very short. The shortening value of a fat varies with a number of conditions and it is therefore absolutely essential in the comparison of two fats that a standard procedure be used. Factors other than the fat that influence the shortening power are (1) manipulation, (2) temperature, (3) ingredients other than flour and fat and their concentration, and (4) concentration of fat. When two fats are compared, all techniques are standardized as carefully as possible. In general, lard has a very high shortening value and butter a lower value, with hydrogenated shortenings intermediate. At least this is true in pastry made from water, flour, and fat. However a fat which ranks high as a shortening agent for pastry is sometimes lower on the list when used in sweet cookie dough.

Several theories to explain why one fat has a greater shortening power than another have been advanced, but none can explain all observations. The structure of doughs and batters is so complex and our knowledge is so incomplete that it is not possible to explain variation in shortening power.

IMPORTANCE IN DIET

Today the quantity of fat and other lipids in a typical American diet is quite high, often accounting for 40 per cent of the calories. This is caused by the relatively high caloric content of one gram of fat—approximately 9 calories (kilocalories) per gram. Although fat and other lipids are present in meat, the amount found in fruits and vegetables is usually rather low. These foods are often enriched with fat in the form of butter or margarine during their preparation. Desserts such as cakes, pastries, and puddings usually have considerable quantities of fat added in their preparation.

The chief contribution of fats and other lipids to the diet is their energy value. In America where at the present time want is seldom known, and where overeating is often a problem, the high contribution of fat may be undesirable. Some nutritionists and physicians now feel that the fat content of the American diet is too high and that for optimum health it must be lowered.

A high fat intake shows a positive correlation with blood cholesterol level in a number of studies. A high blood cholesterol level has been equated with incidence of atherosclerosis although this evidence is by no means conclusive. Unfortunately, it is difficult to measure atherosclerosis in a living man and to determine whether atheromas, the fatty placques in the walls of the blood vessels, are forming, but determination of blood cholesterol is easily run. Thus many experiments have been conducted on blood cholesterol level with the assumption that this measures atherosclerosis.

Much attention has been directed recently to the difference in effect of turated and unsaturated fatty acids. Substituting oils with abundance of saturated fatty acids for hydrogenated fats and saturated fats such as tter and beef tallow often lowers blood cholesterol levels.

Fat and other lipids also have a role in the satiety value of a food. The ling of satisfaction which we derive from a fat, the way it "sticks to r ribs," is the result of numerous physiological reactions, not all of which 2 known. The length of time food stays in the stomach is one reaction iich gives a feeling of satisfaction and delays the onset of hunger. Fat reds the rate at which food leaves the stomach and in this respect increases 2 satiety value of the food. If too much food is eaten at one meal, this ect may prolong the feeling of being "stuffed" and may actually lead to scomfort. Nevertheless, when food is eaten in moderation, one of the luable roles of fat and other lipids is in their satiety value.

Fats and other lipids also contribute essential fatty acids to the diet. hile it has never been unequivocably demonstrated that these fatty acids a essential to man, the work with eczema in babies indicates that we may be a real need for small amounts of linoleic, linolenic, and arachidonic ids in the diet.

Fats are also solvents for the fat-soluble vitamins that are always natilly introduced into the diet in the fatty portion of the food. The fatuble vitamins are vitamins A, D, E, and K, and the provitamins A, the rotenes. All of these vitamins are important to optimum health.

The fats and other lipids are, therefore, important in the diet for a numr of reasons. Whether or not the amount can exceed optimum is yet to be oved.

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HAPTER THREE

Carbohydrates

THAVE HAD an introduction to the chemistry of some of the carbohyates present in foods. You are familiar with the monosaccharides, the xoses—D-glucose, D-galactose, D-mannose, and D-fructose—and perhaps the some of the pentoses—xylose and arabinose. Formulas are given for a common hexoses.

You have studied the disaccharides—sucrose, lactose, and maltose—and the polysaccharides—starches and glycogens. A review of the chemistry of these important compounds can be found in any organic chemistry or biochemistry book.

MONOSACCHARIDES

The pentoses are monosaccharides that contain five carbon atoms. Those that are important in foods, arabinose, xylose, and ribose, are aldoses

α-L-arabinose

rhamnose is a methyl aldose. Like the hexoses, these carbohydrates ring compounds composed of five carbons and one oxygen atom nose) or four carbons and one oxygen (furanose); and it is the ring that occurs in polysaccharides.

α-L-rhamnose

re complexity of the mixture of monosaccharides in plant materials, the sand vegetables of our diet, is not always recognized. The presence e most abundant carbohydrate is sometimes so thoroughly stressed the presence of others is overlooked. Table 3.1 illustrates the large

TABLE	3.1.	SUGAR	C	ONTER	TP	OF	POTA	TOES	G	ROWN	IN	1954	
	AS	DETERM	NE	D BY	P	APER	CHR	OMA	TO	GRAPH	Υ		
	(1	mg/100	g	RAW	PO	TAT	OES),	AV.	8	TESTS			

Sugars	Katahdin	Red Kote	Red Pontiac	Russet F
Maltose	12.0	0.0	51.0	0.0
Sucrose	47.0	66.0	66.0	28.0
Fructose	42.5	70.0	35.5	40.5
Mannose	35.0	0.0	33.5	0.0
Glucose	47.5	45.5	75.0	26.7
Xylose	24.5	42.8	60.0	0.0
Total	208.5	224.3	321.0	95.2

From Habib, A. T. and Brown, H. D., Food Technol. 11, 85-89 (1957).

number of monosaccharides present in a plant part, this time the pota Data are given for the number of milligrams of each monosaccharide part 100 g of tissue for several varieties. The amounts of some are quite low.

DISACCHARIDES

The common disaccharides are already known to you. They are and drides of two monosaccharides (monoses) and include sucrose, lacto maltose and cellobiose. Sucrose is widely distributed in the plant kingdo although sugar cane or sugar beets are the commercial source of mosugar. Maple sugar is also principally sucrose. Lactose occurs in the mof all mammals while maltose and cellobiose occur in low concentration in plants and processed foods.

mercially in many crystal sizes from extremely fine to very coarse. It is often used as a syrup sold as "liquid sugar." This prepared syrup is icularly useful for canners and other processors who use large quantito solutions and is sold in containers of many sizes up to tank cars. Ips are also available in which the sucrose has been partially inverted to ose and fructose. These syrups can be prepared with a higher container of solids since fructose has a very high solubility and glucose not readily crystallize. The graph (Figure 3.1) shows the improvement of ability with increased inversion at various temperatures. Notice that a certain critical concentration of invert sugar is reached, the solubil-leclines. The sweetness of these inverted sugars is comparable to ose. The graph (Figure 3.2) shows a comparison of hexoses with ose. Only fructose is above the equality line. Dulcin is a noncarbohy-sweetner.

Estimers syrups are also available on the market. They contain not only use but also some of the inorganic and organic compounds present in or beets or formed in processing. They vary in color from pale yellow link brown and possess considerable flavor other than the sweetness of use.

that cannot be digested by man and which are, therefore, not of direct that significance. However, many of these compounds are of great it use in the development of the plant and give those plant products

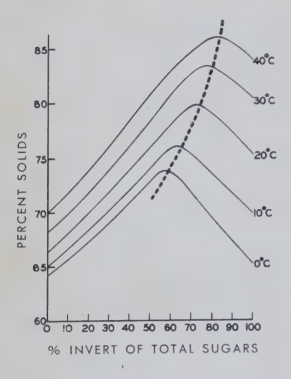


FIGURE 3.1. SOLUBILITY CURVES F SUCROSE-INVERT SUGAR MIXTURES VARIOUS TEMPERATURES. The dotted lishows peak solubility composition for various temperatures. Data of Jur Nelson, and Sherrill. Reproduced from Davis, P. R., and Prince, R. N., "Lique Sugar in the Food Industry," in "Use Sugars and Other Carbohydrates in the Food Industry," Advances in Chem. See 12, American Chemical Society, Washington, D. C. (1955).

used as foods much of their characteristic texture in the fresh and cooke states. These groups of compounds are the celluloses, hemicelluloses, are pectic substances. The lignins may be related to the carbohydrates, but these compounds will not be studied since they are deposited in appreciab amounts only in old, woody plant tissues that are not considered desirab for food.

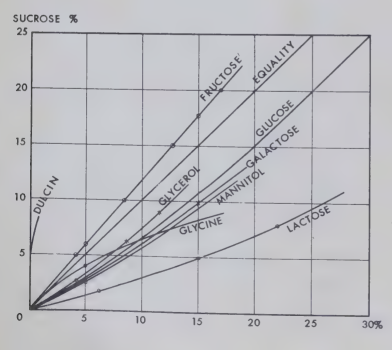
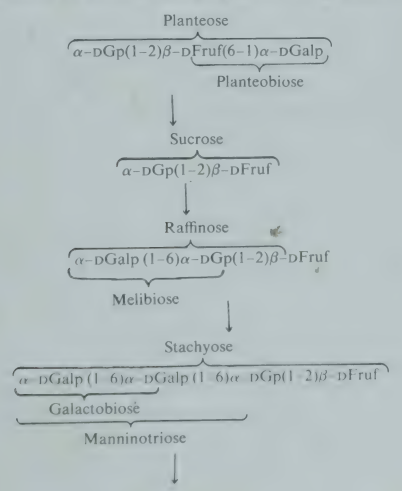


FIGURE 3.2. REL TIVE SWEETNESS VARIOUS COMPOUN COMPARED CROSE FROM DATA CAMERON AND DAH BERG. Reproduced fro Cotton, R. H., Rebe P. A., Maudru, J. and Rorobaugh, ("The Role of Sugar the Food Industry," "Use of Sugars at Other Carbohydrates the Food Industry Advances in Chem. Se 12, American Chemic Society, Washingto D. C. (1955).

GOSACCHARIDES

he oligosaccharides are anhydrides of several monosaccharide residues, as the word "several" is not precise, neither is the prefix "oligo-" ally any compound which contains ten or fewer monosaccharide units assed as an oligosaccharide while those which contain more than ten called polysaccharides. Actually, in nature the most abundant oligocharide is sucrose, with two monosaccharide units and raffinose, with e. All compounds that are well known have less than six monocharide units. An oligosaccharide may be composed of the same monocharide units; however, often it is composed of different units.

he raffinose family is widely distributed and these compounds could lily be produced in large quantities commercially if there were a dead for them. They are found most frequently in seeds, roots, and underand stems. For example, the quantity of raffinose in the seeds of mes is equal to or greater than the quantity of sucrose; and in cotton or soybean meal both raffinose and stachyose occur in abundance. 1ch⁷ shows the relationships between the structures of the individual osaccharides of the raffinose family by the following scheme:



$$\frac{\alpha \ \mathsf{DGalp}(1-6)\alpha - \mathsf{DGalp}(1-6)\alpha - \mathsf{DGalp}(1-6)\alpha - \mathsf{DGp}(1-2)\beta - \mathsf{DFruf}}{\mathsf{Verbascose}}$$

where:

$$G = glucose$$

f = furanose

Fru = fructose p = pyranose

This scheme shows the ether link between the monosaccharide unit conventional means. Thus α or β refers to the configuration of the droxyl group formed in the ring structure of the monosaccharide; and numbers refer to the carbons, numbering in glucose from the aldehyde

bon as 1 to the primary alcohol carbon as 6. The letters f or p refer furanose or pyranose rings. Thus stachyose has a structure as shown:

POLYSACCHARIDES

These compounds are anhydrides of one or more monosaccharides which a large number of units are combined. These high molecular wei compounds are exceedingly difficult to study. Purification is an ardu and tedious task, and then, to demonstrate that a pure compound been isolated is difficult. Even when the presence in a sample of only type of molecule is reasonably certain, determining the molecule's struct

ints problems. However, investigation of the structure of many classes rege molecules (they are called macro molecules) has progressed rapidly g the past two decades and advances in the study of one type of moleculen pushes forward the study of other molecules. Cellulose has been ed intensively; and although it is still impossible to describe with plete assurance a cellulose molecule, much is known about its structure hemicelluloses, gums, and pectic substances are not as well with although many of the details of their structures have been ascerted. We can describe them with some reservations.

any polysaccharides are composed of only one repeating monosacchaand even where a larger number exists they appear to be arranged in a matic fashion. *Homopolysaccharides* possess only one repeating unit a heteropolysaccharides contain more than one. Most polysaccharides composed of aldoses or their derivatives and the hydroxyl group ed in the ring structure on carbon-1 (the anomeric carbon) reacts to the ether link that holds the monosaccharide residues together. This paxyl group may react with any other available hydroxyl on the adat monosaccharide, except the one on carbon-1, to form the polysacde chain. Usually the one with which it reacts is the same throughhe chain.

ou are familiar with the structure of the two types of starches: amyloses long chains of glucose residues and the amylopectins with branching as that form a "bushy" molecule. You will remember that the glycoare similar in structure to the amylopectins, although the number of se residues in the chain is smaller. (These can be reviewed in any nic chemistry textbook.)

Jose

lulose is the compound that gives strength to plant tissues. It is ded in the cell walls of all seed-bearing plants in varying concentration, orms long fibers that strengthen stems and ribs of leaves. On acid hysis of cellulose, glucose is formed in yields of 95 to 96 per cent. This dictes that the structural unit in a cellulose molecule is glucose. Callulose is treated with an acetylating agent, it forms cellobiose at 2, the only disaccharide produced. Cellobiose is a disaccharide of cose units which differs from maltose in the configuration of the 1-m. In maltose the alpha configuration occurs, while in cellobiose the ite configuration, beta, exists. We can conclude from the formation ellobiose derivative that the beta configuration exists in cellulose, studies have shown that the link between the glucose residues is a 1-4 lie same one that is shown in cellobiose. The exact number of glucose

residues in the chain is still unknown, but it is certain that the number must be large. Attempts to determine the molecular weight of cellulose is lated from various plants, by viscosity measurements, by the ultracentrifuge, and by end group analysis have not given consistent result Most determinations give ranges of values and indicate that the samples a composed of molecules of different molecular size.

Schulz and Marx²⁴ have found that the degree of polymerization of cel lose from fibers in cotton, cotton linters, ramie, china grass, and flax between 6500 to 9000 glucose units, while those from the first leaf past to cotyledon on plants, from root tips, or bacteria show considerable vartion and are considerably lower in molecular weight. These celluloses a probably representative of those found in foods.

X-ray studies of cellulose fibers indicate that the fibers contain areas crystallization and areas of disorganization called amorphous cellulo. These data indicate that in the crystalline areas the cellulose moleculare oriented parallel to the fiber axis. The dimensions of the cells have been calculated, and it is believed that one cellulose molecule lies in ordirection, while the neighboring molecule lies in the opposite direction. To molecules are so close together that they probably are held by hydrog bonds between oxygen atoms. In the amorphous portions of the fiber much looser arrangement exists; in this part adsorption of water and swe ing readily occurs. Also this part of the fiber is believed to be flexible a capable of distortion without breaking. If this hypothesis is true, the property makes a cellulose fiber of peculiar value to the plant, strengthese



FIGURE 3.3. CELLU-DSE FIBER IN CELL WALL F Valonia LIE IN AL-ERNATE LAYERS. Electron icrograph × 15,000.

Courtesy of R. D. Preston.



FIGURE 3.4. CELLU-DSE FIBERS ON THE NER SIDE OF THE CELL ALL OF Valonia. Granur material may be protoasm. Electron microaph × 22,500.

Courtesy of R. D. Preston.

g a stem or leaf and yet not rendering it rigid and susceptible to break-

tarch

Starch is the reserve carbohydrate of plants and occurs as granules in cell in plastids, separated from the cytoplasm. Under the microscope plastids in some plant tissues can be seen filled with granules and nen the cells are ruptured, the granules stream out. The size and shape of arch granules is characteristic of their origin, and anyone trained can adily identify the origin of starch granules with a microscope.

Most starch granules contain both amylose and amylopectin molecules.

v or glutmous starch from corn and other cereals contains little or no

amylose, while a sugary mutant corn and some of the legumes contain amylose in greater abundance than amylopectin. Amylopectin is usuall the more plentiful type of starch. (See Table 3.2.)

Amylose molecules are straight chain polysaccharides in which α -D-glucose units are joined 1–4. Chain lengths vary from 250 to about 350 glucose units, and the long molecules appear to be coiled in an α helix Amylopectin molecules are branched at carbon 6 on the glucose unit to form a 1–6 ether link. The length of the linear units in amylopectin in only 25 to 30 units but molecular weights show that 1000 or more glucose units are combined in a molecule.

The ability of a natural starch to form pastes and gels differs considerably with the source of the starch. Amyloses are believed to form gel more readily because the linear shape allows the formation of a three dimensional network with ease. However, these molecules associate and crystallize readily and the starch dispersion undergoes retrogradation (see p. 107). Amylopectin molecules, with their bushy structure, do not crystal lize readily. Waxy starches do not show retrogradation.

Enzymes and Starch. Three types of enzymes, widely distributed in nature, react with starches: α -amylases, β -amylases, and phosphorylases. The amylases are particularly interesting to the food chemist since they are often used to modify starches.

Alpha Amylase. Alpha amylases are also called "liquefying" or "dextrogenic" amylases from their action on starches. They occur widely distributed in the plant world often associated with β -amylases. Dormant, ungerminated seeds sometimes possess α -amylases in small quantities, but on germination all of these seeds rapidly develop these enzymes. Thus alpha amylases can be readily prepared from sprouting grains. Alpha



FIGURE 3.5. MICELLAR ORGANIZATION WITHIN SWOLLEN STARCH
GRANULE, ACCORDING TO K. H.
MEYER. Reproduced from Schoch,
T. J. and Elder, A. L., "Starches
in the Food Industry," in "Use of
Sugars and Other Carbohydrates
in the Food Industry," Advances in
Chem. Ser., 12, American Chemical Society, Washington, D. C.
(1955).

nylases also occur in saliva and pancreatic juice and are produced by a umber of bacteria and fungi.

Alpha amylase catalyzes the hydrolysis of starches into low molecular right dextrins with great rapidity. (See Figure 3.6.) In a short time a trch dispersion liquefies as the molecular weight of the colloid is decased and soon the solution is filled with dextrins of approximately 6 acose units along with a small amount of maltose. The bonds hydrolyzed e the 1-4 ether links. The 1-6 links in amylopectic molecules are by-tssed. Any phosphate esters are likewise left intact.

Beta Amylase. This enzyme is also widely distributed. Beta amylase is lled the "saccharifying amylase" because the chief product of its hyolytic catalysis is maltose. (See Figure 3.7.) A relatively pure extract of amylase can be made from germinating soy beans. "Diastase," a comercial extract from barley malt which is widely used in industry both for od and many other products, is a mixture of alpha and beta amylases.

Beta amylase forms primarily maltose and the reaction will go almost to impletion. The 1 6 link appears to be somewhat sensitive to hydrolysis the presence of β -amylase as well as the 1 4 link between glucose its. The barrier link that prevents the formation of 100° maltose is obably the β -link which occurs infrequently in amylose and amylopectin the 1-carbon of glucose.

Technology of Starch. Starch is produced in large quantities in the nited States from a number of plant sources. Cornstarch is the most im-

TABLE 3.2. AMYLOSE CONTENT OF SOME STARCHES*

	S. Jack M.	Amylose Content, %	
	Sugary Mutant Corn	70	
	Steadfast Pea	67	
	Alderman Pea	65	
	Chick Pea	33	
	Buckwheat	28	
	Barley	27 •	
	Sorghum	27	
	Commercial Corn	26	
	Wheat	25	
	White Potato	23	
	Arrowroot	21	
	Sweet Potato	20	
	Tapioca	18	
	Waxy Barley	3	
	Waxy Corn	0-6	

Fr in Whistler P D and Smart C 1, "Polysaccharide Chemistry" p. 242. Academic Press. Inc., York, N. Y., 1953.

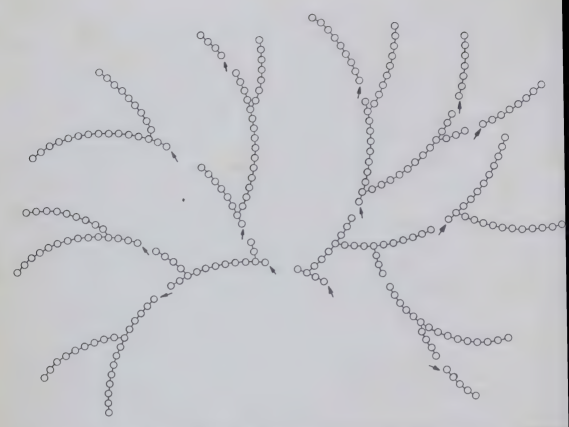


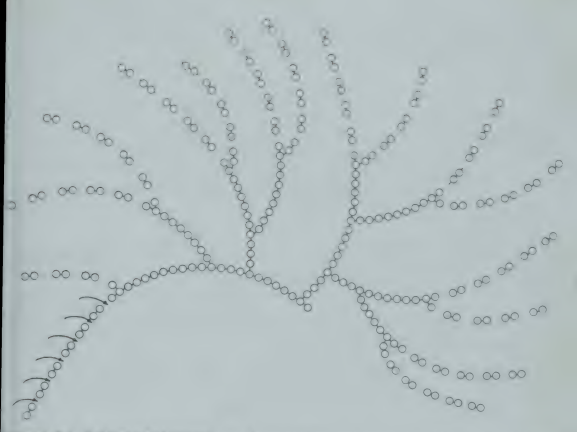
FIGURE 3.6. α -Amylase Activity on Amylopectin with the Formation of Dextrin of Low Molecular Weight. *Advances in Enzymol.*, 12, 390 (1951).

Courtesy of P. Bernfeld

portant. The most common method is the one described below but many modifications exist.

The corn is steeped for 35 to 45 hr in water at 50°F with sufficient sulfur dioxide added (0.15 to 0.2 per cent) to prevent growth of microor ganisms and perhaps start the disintegration of protein. During steeping the corn picks up water and is then readily ground to separate hulls and germ. The starch, protein, and water-soluble substances form a slurry that is separated from the germ. The slurry is then finely ground and separated from hulls and coarse particles. More sulfur dioxide is then added and the temperature increased to within the range of 85°F to 90°F to make the separation of the starch granules from the protein accoust material easier. The granules are separated by various methods such as centrifugation of filtering and are washed and dried. The starch is sold in several forms a Pearl is rough-ground from the dryers while powder is finely ground and even sifted. Lump or crystal is the other form available on the market.

Many modified starches are on the market but those most commonlused in the food industry are the thin boiling types. They are the result of partial hydrolysis of the starch, usually by sulfuric acid or some other



ICURE 3.7. β-AMYLASE ACTIVITY ON AMYLOPECTIN WITH THE FORMATION OF MALTOSE A HIGH MOLECULAR LIMIT DEXIRIN. Advances in Enzymol., 12, 394 (1951).

Courtesy of P. Bernfeld.

d. Before drying the starch slurry is treated with 0.1N H₂SO₄ at 50°C for o 24 hr. Hydrolysis of some of the bonds in the starch results in a oduct that will disperse in boiling water to yield a dispersion with a cosity not much greater than water. The more extensive the modification the starch, the greater the effect on lowering viscosity.

uctans

Fructans are polysaccharides composed of fructose residues and are desited in the roots and tubers of a number of plants as well as in the ves of some grasses. A number have been isolated and the structure of ne is fairly well known. Some fructans are linear while others are nebed, with either a 2-1 or 2-6 link between the fructose residues, and a fairly low molecular weight. Since they are easily isolated and since cose is the sweetest of the hexoses, fructose syrup has been produced numercially.

annans and Galactans

these as well as the heteropolysaccharides of galactose and mannose, a lactomannans, have been found in plant and animal tissues. These last

are used as thickners and are produced commercially from seeds. They a briefly discussed under gums.

Chitin

Chitin is a homopolysaccharide that occurs as a structural compound lobster, shrimp, and crab shells as well as in some insects and fungi. To shells contain calcium carbonate, proteins, and other compounds, but the polysaccharide is mainly responsible for the rigidity of the tissue. It easily extracted. The repeating unit in chitin is 2-N-acetyl glucosamin linked β -1-4.

Portion of Chitin

Hyaluronic Acid and the Chondroitin Sulfates

The heteropolysaccharides of animal tissues have been studied for many years and several of them in the past decade have been well characterized. The problem of separation is a difficult one and there is still a great decof confusion in this field in nomenclature and definition. Attempts in the past to determine structure with crude mixtures rather than with pure compounds have added to this confusion. Hyaluronic acid and the chondroitis sulfates are now well known and the structures at least partially decade.

Portion of Hyaluronic Acid

termined. Both occur in the ground substance of connective tissue. Hy aluronic acid is more abundant in skin and soft tissues while the chondro tin sulfates are more abundant in cartilage. Hyaluronic acid can be iso ed from umbilical cord and chondroitin sulfates from bovine nasal tum. Probably in connective tissue these carbohydrates occur linked to otein.

Iyaluronic acid appears to be a linear molecule or one in which there ittle branching of alternate units of 2N-acetyl glucosamine and gluonic acid. These units are joined $\beta = 1/3$ and $\beta = 1/4$. Some workers find a hyaluronic acid is difficult to methylate and believe this indicates a ratified structure for the molecule rather than a linear one. Viscosity studindicate a molecular weight of 200,000 to 400,000.

Hyaluronic acid is found widely distributed. All connective tissue apars to contain it or derivatives and its jellylike quality when dispersed in ter gives these tissues their particular softness and elasticity. Vitreous diaqueous humor, synovial fluid, and pleural fluid all contain it. In these ids its rather high viscosity make it an excellent lubricant.

Three chondroitin sulfates have been isolated and are designated A, B, 1 C. Chondroitin sulfate A and C are polysaccharides composed of equal less of 2N-acetyl galactosamine, p-glucuronic acid, and sulfuric acid, ey are joined through beta 1-3 and 1-4 links.

Portion of Chondroitin A and C Chain

A olecular weights as high as 2,000,000 have been found for chondroitin ate A, but chondroitin sulfate C is smaller in size. Chondroitin B is eved to differ from A and C by possessing 1-iduronic acid in place of

α-L-iduronic Acid

D-glucuronic acid. This differs from glucuronic acid in the configuration carbon 5.

Chondroitin sulfates A, B, and C differ in their optical activity and the resistance to enzymatic digestion by either testicular hyaluronidase or the hyaluronidase of pneumococci. Chondroitin sulfuric acid has been isolated from cartilage with good yields: pig nasal septum contains 41 per ce while the cartilage from the epiphysis of an infant yielded 20 per cent.

TABLE 3.3. DISTRIBUTION OF HYALURONIC ACID AND CHRONDROITIN SULFURIC ACID A, B, AND C*

Group	Tissue	Hyaluronic Acid	ChS A	ChS B	Ch
I.	Vitreous Humor	+	10000	_	_
	Synovial Humor	+	-		_
	Mesothelioma	+	_	-	_
II.	Hyaline Cartilage	Anna	+		+
III.	Heart Valves		_	+	+
	Tendon (Pig and Calf)	±	_	+	+
	Aorta	-	_	+	+
IV.	Skin (Pig and Calf)	+	_	+	_
	Umbilical Cord	+	*****	_	+

^{*}From Meyer, K. and Rapport, M. M., Science, 113, 596-599 (1951).

Meyer and Rapport¹⁷ give the distribution of hyaluronic acid ar chondroitin sulfuric acid A, B, and C from ground substance polysacch rides in the tissues studied so far, as shown in Table 3.3. Chondroitin su furic acids are believed to be weakly linked to protein in connective tissues since they are readily extracted with solutions of calcium chloride.

Mucoitin sulfuric acid (hyalurono sulfuric acid) is a heteropolysacch ride which has been less extensively studied than chondroitin sulfuric acid or hyaluronic acid. It is closely related to them since on hydrolysis it form glucuronic acid, 2-desoxy-2-amino D-glucose (glucosamine), acetic acid and sulfuric acid. It has been isolated from cattle corneas and gastr mucosa. The compound from corneas is readily attacked by the enzyme hyaluronidase and is therefore believed to be the mono-sulfuric acid ester of hyaluronic acid. This is not true of the mucoitin sulfuric acid isolated from the gastric mucosa. It resists attack by hyaluronidase and must therefore possess a different structure.

Mucopolysaccharides and mucoproteins are complex compounds relate to carbohydrates and are of considerable importance in mammals. The nomenclature is not yet precise, but the groups of compounds are usuall classified either as mucopolysaccharides or mucoproteins on the basis of the amount of carbohydrate or protein present, although no exact dividin e can be given. These mucosubstances are slimy compounds that act as vicants, protect epithelial surfaces, or form part of the cell wall and e many other functions.

The carbohydrate portion of these molecules is composed of hexosme, monoses, and often sialic acid. The hexosamines are either glucosme or galactosamine while the monose may be galactose, mannose or ose or combinations of them. In bacterial cell walls the pentoses, mnose and arabinose, occur. The carbohydrate portion is often large in some mucoproteins appears quite small as a prosthetic group. All se polysaccharides are bound tightly to protein. This is a different kind bond than that which occurs between the chondroitin sulfates or hyronic acid and protein.

ialic acid is the name now given to all acylated derivatives of neuraic acid. The one most commonly encountered appears to be N-acetyl raminic acid. Neuraminic acid is now considered to be the condensate nannosamine and pyruvic acid.

Sialic Acid (N-acetyl Neuraminic Acid)

1 any of these polysaccharides are present in small amounts in meat and t products.

nicellulose

ant cell walls are readily visible under a microscope because of thickand the frequent occurrence of numerous layers. The most abundant
pound present is a cellulose. Long fibers are embedded in a matrix
menting compounds and in older tissues which have become woody
nerasted with lignins. In analysis of plant tissue and attempts to
ate the compounds that make up the cell wall, lignins are first reed This leaves "holocellulose," which appears to be an intimate mixof cellulose and other compounds which are soluble in aqueous alkali.
e alkali-soluble substances are called hemicelluloses and are not well
ed No satisfactory classification of the hemicelluloses has been posvet although some generalizations are given.

It is generally agreed that the following differences exist between ce lose and hemicellulose molecules:

(1) Celluloses have a higher degree of polymerization (more monos charide units) per molecule than hemicelluloses.

(2) Celluloses are less soluble in alkali and less readily hydroly by dilute acids than hemicelluloses.

(3) Celluloses are fibrous, while hemicelluloses are nonfibrous.

(4) Celluloses yield D-glucose on hydrolysis, while hemicelluloses yield predominantly D-xylose and other monosaccharides.

(5) Celluloses have a higher ignition temperature than hemicellulos. These criteria are only general, qualitative means of differentiation.

Hemicelluloses are widely distributed in nature, but research has detered for the most part on fractions isolated from wood. Most hemice lose is composed primarily of the pentose, xylose, often combined with methyl uronic acid, although hemicellulose fractions from conifers continuannose, one of the hexoses, as the principal monosaccharide residentable 3.4 shows that many other monosaccharides, both hexoses a pentoses, have been isolated from hemicellulose fractions. Plant hemicelluses other than wood which have been studied have a similar composite with xylose the principal monosaccharide residue. This is shown in the din Table 3.4. The uronic acids found have not been well characterized, though some have been identified as glucuronic or galacturonic acids, their methyl ethers. Hirst¹¹ summarizes our knowledge of the structure of few fairly well defined hemicelluloses in schemes which represent polysicharides composed of a number of monosaccharide units. See Table 3. In this table where the number is known, it is given.

TABLE 3.4 COMPOSITION OF HEMICELLULOSES*

Nylose Methyluronic Glucose Galactose Mannesc Arabitose Rhan	s	Pentoses			Hexoses				
wood mi A	- Glu-	Rham-		Mannose		Glucose		Vylose	(
e birch	curone	noxe				23% in sap-	17%	82% 60%	mi A mi B k locust
Invite						not neart-	7.8-19.5%		e birch
spruce \(\sqrt{ \sq}} \sqrt{ \sqrt{ \sqrt{ \sqrt{ \sq \sqrt{ \sq							1 mole		
Spruce V V V V V V V V V	sm.	sm	sm.	sm.	sm.	sm.		chief	nwood
e Pine 44-50 10.5-15 sm. 36-46% /ood s sol. 7.7% 3.3% chief Pine act. A act. B act. C spruce A B C fa Hay chief some r Cane 22 moles 0.9 moles per (no me.) stalk 19 moles 2 (no me.) 7	SIII. √			√	V	√		\checkmark	spruce
s sol. 7.7% 3.3% chief Pine act. A act. B act. C 45.2 spruce A B C 44.9 fa Hay chief some r Cane 22 moles 0.9 moles cer (no me.) stalk 19 moles 2 (no me.) 7	v	v	v	36-46%		sm.	10.5-15	44-50	
Pine act. A act. B act. B act. C 45.2 spruce A B C 44.9 fa Hay chief some r Cane 22 moles 0.9 moles oer (no me.) stalk 19 moles 2 (no me.) 7				chief			3.3%	7.7%	
act. B act. C act. C 45.2 spruce A B C 44.9 fa Hay chief some sm. r Cane 22 moles 0.9 moles (no me.) stalk 19 moles 2 (no me.) 7									
act. C spruce A B C fa Hay chief some r Cane 22 moles 0.9 moles (no me.) stalk 19 moles 2 (no me.) 45.2 9.3 26.4 44.9 5m. 1 mole									
A B C C A B									act. C
B C C 44.9 fa Hay chief some sm. r Cane 22 moles 0.9 moles no me.) stalk 19 moles 2 (no me.) 7									spruce
C 44.9 fa Hay chief some sm. r Cane 22 moles 0.9 moles no me.) stalk 19 moles 2 (no me.) 7									
fa Hay chief some sm. T Cane 22 moles 0.9 moles 1 mole (no me.) stalk 19 moles 2 (no me.) 7								٠	
r Cane 22 moles 0.9 moles 1 mole oer (no me.) 7				44.9				-1-:-C	
stalk 19 moles 2 (no me.) 7			sm.						
			1 mole					22 moles	
fulls $\sqrt{1 \text{ (no me.)}}$ 2 mol $$			7				2 (no me.)	19 moles	stalk
			√		2 mol		1 (no me.)	√	Iulls
rto 18-20 1			1					18-20	
maple uronic methoxy									
ct. IA 46.1 17.1 2.7									
IB 48.7 15.8 2.3									
II 54.7 22.8 2.6									
III 79.2 12.2 2.1 IV 80.9 9.3 2.3									

off W & L E and John E C , "Wood Chemistry." 2nd ed , Chap 10, "The Hemicelluloses." old Publishing Corp., New York, N. Y. (1952).

TABLE 3.5. STRUCTURE OF HEMICELLULOSES*

Source	Structure	No. of Monosacche Units
Esparto grass	X14X14X14X 3 X14X1	$\xrightarrow{x y} x + y + z =$
Corn cob xylan	similar	2/ 32 1 9 1 2 -
Wheat flour	X14X14X14X14X 3 3 2 -1A 1A1A	
Esparto grass	X14X14X14X 3 3 1A 1X41X	
Wheat straw	X14X14X14X14X14X 3 3 1A 1GA	50 <i>x</i>
Oat straw	X14X14X14X14X 3 2 1A c1GA	
Pear fruit	$X1 \dots 4X1 \dots 4X1 \dots 4X1 \dots 4X1 \dots 4X$ $X1 \dots 4X1 \dots 4X1 \dots 1GA$	$\frac{xy}{z/GA} x + y + z =$
Beechwood	X14X14X14X 2 1MGA	
Algal xylan Phododymeia paln (Red seaweed)	X14X13X1X	Ratio 1:12:3
Barley gum	$\dots 4G\beta 1-3G\beta 1\dots$	
Larchwood	$Gal\beta 1$ -6 $Gal\beta 1$	
	1β Gal6	
Guar	$4M\beta 14M\beta 1 \dots$	
where:	$\dots 1\alpha$ Gal	
M = Mannopy	ranose Dyranose Opyranose Ayranose Opyranose O	

^{*}From Hirst, E. L., J. Chem. Soc., 2974–2984 (1955).

st's scheme attempts to show not only the branching of the molecules the carbons at which bonds are established. Thus wheat flour hemiclose has a "back bone" of a chain of xylose residues, joined 1.4 with aching by arabinose moieties on either carbon 3 or occasionally carbon he formulas show this. (See the formula presented above on p. 84.) The aer low molecular weights are demonstrated by rather low number of nose residues in esparto grass, wheat straw, pear fruit hemicellulose (80, and 120).

:tic Substances

hese consist of a group of carbohydrates which are of considerable inst in foods of plant origin. They comprise not only those substances atins) in fruits and vegetables which are capable of forming gels with ar and acids but a number of other compounds as well. The exact loon of these compounds in a tissue is still a matter of dispute since methods of chemical morphology are not yet specific enough. Most kers are agreed that pectic substances occur in the middle lamella been cells and perhaps in the cell wall, but whether they are ever coned within the cell is questionable. The structure of the pectic subscess has been the object of a great deal of research and considerable wledge has accumulated. The chief products of the hydrolysis of pectic stances are galacturonic acid, a derivative of galactose in which the 6-bon is oxidized to a carboxyl group, and methyl alcohol.

s indicates that pectic substances are polysaccharides of galacturonic or of its methyl ester. Whether other groups are present in the moless difficult to determine because pectic substances are hard to purify, some apparently pure samples of pectic substances have been pred which contain only galacturonic acid or its methyl ester, when other eties are found among the hydrolysis products, it is difficult to decide ther they are contaminants or intrinsic parts of the original molecules, he pectic substances have been studied for many years and the nomentar in the early days became very confused. In 1926 a committee of the

American Chemical Society attempted to standardize this nomenclated Its report was revised and adopted (see Kertesz¹⁴) as official in 1943. The definitions are given here:

Pectic substances "Pectic substances" is a group designation for those complex collecarbohydrate derivatives which occur in, or are prepared from, plants and contain a proportion of anhydrogalacturonic acid units which are thought to exist in a chainlike obination. The carboxyl groups of polygalacturonic acids may be partly esterified by megroups and partly or completely neutralized by one or more bases.

Protopectin. The term protopectin is applied to the water-insoluble parent pectic subst which occurs in plants and which, upon restricted hydrolysis, yields pectinic acids.

Pectinic acids. The term pectinic acids is used for colloidal polygalacturonic acids taining more than a negligible proportion of methyl ester groups. Pectinic acids, usuitable conditions, are capable of forming gels (jellies) with sugar and acid or, if ably low in methoxyl content, with certain metallic ions. The salts of pectinic acids either normal or acid pectinates.

Pectin. The general term pectin (or pectins) designates those water-soluble pectinic a of varying methyl ester content and degree of neutralization which are capable of form gels with sugar and acid under suitable conditions.

Pectic acid. The term pectic acid is applied to pectic substances mostly composed colloidal polygalacturonic acids and essentially free from methyl ester groups. The of pectic acid are either normal or acid pectates.

The molecular weights of various pectic substances extracted from pl material have been determined by osmotic methods, the ultracentrifu end group analysis, and by viscosity, but no agreement is apparent as y This is not surprising since separation of individual molecular species in pure state is very difficult, and the pectic substances present in plants probably complicated mixtures. End group analysis indicates a range molecular weights from 2500 to 7500 while measurements of pectic si stance derivatives (acetyl and nitropectins) by osmotic pressures give n lecular weights from 30,000 to 100,000. By use of the ultracentrifuge cru extracts give a range from 16,000 to 50,000, although some scientists has found for highly purified products a variation from 33,000 to 117,000. \ cosity measurements give results from 27,000 to 115,000 with one sample high as 280,000. Probably pectic substances occur in plant tissues as a m ture of molecules with a variety of molecular weights. At the moment i impossible to estimate the number of galacturonic acid residues present any one type of pectic substance.

There have been numerous structures proposed for the pectic substant but today the most popular hypothesis is that galacturonic acid resid are joined in linear units by α 1-4 links like those which occur in amylos Pectic acids are the polygalacturonic acids which are essentially free methyl ester groups, while pectinic acids contain some methyl ester groups

ertesz¹⁴ shows the difference between them in the following scheme:

ne 1 4 link is an α ether link between the first carbon on one galacturonic sidue and the fourth carbon on another.

Pectic Substance Chain

be 90 to the plane of the galacturonic acid ring. This differs from it for other polysaccharides and gives the linear unit a screwlike consuration. The number of galacturonic acid residues in each linear unit is 1 hown. Since pectic acids can be prepared from pectinic acids by hypolysis, it is assumed that these two groups of compounds have essentially same structure.

The free carboxyl groups in both pectic and pectinic acids will react metallic ions to form salts. If the acid is completely neutralized, that all carboxy groups react, the product is a normal pectate or pectinat only part of them react, the product is an acid pectate or pectinate. No of the pectic substances exist in plants as acid calcium or magnesium sa

Both pectic acids and pectinic acids are predominantly linear. Howemany workers in the field believe that the screw-shaped linear units of lacturonic acid residues are associated as macromolecules, either throsome sort of branching similar to that which occurs in starches and gligens or by lamination or bundle formation. Three possible types of I which hold the linear units have been suggested: (1) anhydride format between the carboxyl groups, (2) ester formation between carboxyl and coholic hydroxyl, and (3) hydrogen bonding.

(1) Some evidence points to the participation of the carboxyl grouplinks between linear units of galacturonic acid residues. For one thing, equivalent weight is not high enough to indicate one carboxyl for eglucuronic acid residue. Also pectic substances are especially vulnerable alkaline degradation, a characteristic of acid anhydrides. The anhydlink is the following:

$$GC \longrightarrow OH + G \longrightarrow COH \longrightarrow GC \longrightarrow O \longrightarrow G + H_2O$$

(2) The link may be ester in nature although this group is more diffito hydrolyze.

$$\begin{array}{c|c} O & & O \\ GCOH + GOH \rightarrow G-C-O-G \\ \end{array}$$

Either link (1) or (2) would form branched or laminated molecules veconsiderable stability since these are primary valence bonds.

(3) The hydrogen bond which weakly holds many macromolecules gether and is readily disrupted by warming may hold linear units of ga turonic acid residues in laminated molecules. Hydrogen bonding can obsetween hydroxyl groups or between hydroxyl and carboxyl. This type macromolecule will dissociate readily into linear molecules.

Protopectin. Protopectin has been assumed for many years to exist as insoluble molecule in plant tissues, but its exact nature is far from solv. There is always a great problem in separating a large insoluble molecular from the complex mixtures which occur in plants, without producin change in composition. Attempts to study it by staining techniques

wise not progressed far enough to give any information that is not pered with doubt.

'any investigators in the past assumed that protopectin is a large inible molecule formed by the reaction of pectic substances with celluCellulose occurs in the cell walls of plants and protopectin is beed to be in these walls as well as in the middle lamella between the
walls. It is certainly possible that such a combination exists, and such
uge molecule would be insoluble.

Other workers have suggested that protopectin is insoluble because it sts as a calcium-magnesium salt. Protopectin is rendered soluble by hydrosis with acid and they have regarded this action as the substitution of frogen for the calcium and magnesium ions. In support of this hypothethere are also the observations that isolated pectic and pectinic acids of ter high molecular weight form insoluble salts with calcium and magium ions. In other words, these workers consider protopectin as tum-magnesium pectinate. Finally there are some researchers who contributed in a macromolecule built up of numerous pectinic acids and whave speculated on the type of linkages which hold the pectinic acids ether. It may be that ether or ester links form unusually large branched aminated molecules similar to those described for pectic and pectinic is. It is also possible that the pectinic acids are held together by tium ion bridges. Since the calcium ion is divalent, it is possible that it at with a carboxy' group on each of two pectinic acids.

he reactions of pectic substances are very evident in cooking fruits and ticularly in making jelly. When fruits rich in pectic substances are boiled tydrolyze the protopectin they form a juice which will gel when sufficient and acid is added. Overripe fruit in which the pectinic acids have extensively hydrolyzed to pectic acids, make very weak jellies or none at Boiling the juice for long periods often produces syrupy gels for the ne reason. The physical-chemical problems encountered in preparing tir gels are complicated; and although the companies which make compare at pectin either from apple pomace or from citrus fruits have standard their products and have established procedures which insure suc-

cessful jellies, many of the principles behind these procedures are obscure.

Pectin Gels. The pectin which is present in most fruits has been used many years to produce sugar-acid gels. Some fruits such as apple and quare widely known to have excellent jellying power. The pectic substantial which produce firm jellies are relatively high molecular weight mole with a relatively high per cent of methyl ester groups and consequent low per cent of free carboxyl. When a hot aqueous mixture of sugar, and pectins is cooled, it sets into a gel. The sequence of events and effects of ingredients is complex. Several theories have been proposed to plain the gel formation. The factors which influence gelling will be cussed, but their significance will be omitted.

A firm gel depends on (1) per cent pectin, (2) molecular weight of pectins, (3) per cent methylation, (4) per cent sugar, and (5) pH. A dealed ble jelly must be firm enough to stand without appreciable deformated and yet tender enough to spread readily on bread. As the percentage pectins increases in mixtures, the firmness of the jellies produced on coing increases. A satisfactory jelly is produced with approximately 1 per opectin. The amount varies with the quality of the pectin preparation, average molecular weights of the molecules present, and the degree methylation.

The molecular weight of pectic substances is important in determining jellying power. Molecules must be colloidally dispersed before a gel is passible. Thus the protopectins cannot act in gelation until they have be changed to pectinic acids. As a gel forms, the molecules develop a the dimensional network which traps solution in the interstices. If the macules are too short, the network is not continuous in many spots and the is runny or soft.

The methyl ester content of the pectinic acid is another factor which fluences the jellying power of the product. Excellent jellies can be prepared from pectins with a wide range of methyl content, but maximum jelly appears at about 8 per cent. This represents esterification of half of carboxyl groups.

The effect of pH on jellying has been recognized for many years. In home preparation of jellies it is well known that dead-ripe fruit with lower acid content does not yield as good a jelly as partially ripe fruit. Me pectic products do not form a jelly until the pH is lowered to 3.5, and firmness of the jelly increases as the pH is lowered. With very low pH's, amount of pectin can be decreased and a satisfactory gel still formed. Optimum pH is usually found; and if the pH is lowered below this pothe firmness of the jelly diminishes and excessive syneresis develops.

rigar (mainly sucrose) is necessary for the formation of the pectin gels must be present in a minimum concentration. Most jellies are made approximately 65 per cent sugar. If the amount is increased much e this point, crystallization tends to occur on the surface of the jelly occasionally even within the jelly. We find that the same methods h prevent or diminish crystal growth of sucrose in sugar cookery are tive in fruit jellies. Cooking the sugar with acid fosters hydrolysis of ose to form glucose and fructose (invert sugar), or addition of corn p or other glucose or fructose products in small amounts decreases endency to crystallization.

value but it is difficult to measure. A number of methods have been ested to evaluate the ability of a pectin product to form a jelly. Before anufacturer buys a product, he wants to know how much jelly can be e from 1 lb of the material. Jelly grade is defined in terms of the unt of sugar which 1 lb of the pectin will "carry" or gel. If a pound of in will carry 100 lb of sugar and form a satisfactory jelly, it is classed 30 grade pectin. One of the most difficult points is the definition and

surement of "a satisfactory jelly."

riting Time. The time which elapses between the addition of all coments and formation of a gel, called the "setting time," often has a ked effect on the quality of the final jelly. If a gel sets too rapidly bepouring is complete, the jelly never achieves the firmness possible with setting. When setting is very rapid, the mixture is said to "curdle," as I lumps of gel are formed in the pot. A curdled mix is difficult to pour does not fill evenly. With jams and preserves, however, slow setting we the chunks of fruit to settle rather than remain evenly distributed ugh the jam. Commercial pectins are available either as "rapid set" or slow set."

hot mixture does not set until it begins to cool. The rate of cooling all of the physical factors which effect rate of cooling, such as the pot and shape, have an effect on setting time. But when this is controlled, rent pectin products still show a difference in the setting time. A id set" begins at about 88° C (190° F), while a "slow set" forms a jelly 54° C (130° F). A number of patents have been issued for methods atment of pectin products to prolong setting time. They are methods by acid or enzyme treatment alter the pectinic acids by partial hypers so that both molecular weights and total methyl are slightly ged.

ne pH of the mixture has an influence on setting time. Setting is more it it lower pH's. Salts tend to delay setting; and some, such as

disodium phosphate, are regularly added to "slow set" pectins. Buffe decrease the ionization of the organic acids present in fruit juices a this action raise the pH and prolong setting time.

Adjustment of the pH of a fruit juice or mixture is allowed b USDA. If the fruit has too high a pH, it is permissible to add acid. and lactic acids are the most common ones used since they are believed impart good flavor; but phosphoric, tartaric, and malic are also used. acids are either added at the end of the cooking time so that too much drolysis of the sugar is prevented, or in jellies they are sometimes place the containers and mixed through the hot product immediately after ing. To raise the pH of a fruit which is unusually acid, buffer salts are The USDA allows as much as 3 oz of either sodium citrate or sodium tassium tartrate per 100 lb of sugar.

Gel Formation with Low Ester Pectins. Low ester pectins which he methyl content below 7 per cent are able to form gels in the present small amounts of divalent ions, even though the per cent of solids is low. Commercial products are now on the market and are used for types of gels than fruit jellies or jams. They are produced by de-ester tion with either acid, enzymes, or alkali. The products formed from three methods are evidently not identical since they differ sharply in physical characteristics and particularly in their reaction to divalent They are often called "low methoxyl pectins."

Low ester pectins do not require the presence of sugar for the form of a gel as do the high ester pectins. The significance of the require for divalent ions probably lies in the ability of the ions to react with carboxyl group on two molecules of pectic acid and form a bridge bet them.

The low ester pectins are used in the preparation of low solid salad desserts. Tomato aspic, for example, can be formed with low ester petut will not gel with the high ester pectins. They can also be added to before freezing to reduce the amount of "run-off" when the fruit is that

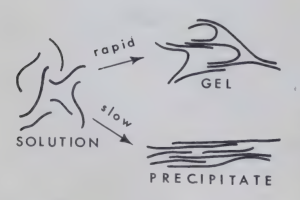


FIGURE 3.8. MECHANISM OF R GRADATION OF LINEAR FRACTION STARCH. Reproduced from Schoch, and Elder, A. L., "Starches in the Industry," in "Use of Sugars and Carbohydrates in the Food Industry, vances in Chem. Ser., 12, American Cal Society, Washington, D. C. (1955).

ing for many foods is possible with a bath of low ester pectins to which all amount of calcium chloride is added. The coating can be sprayed calcium chloride to "tan" the surface. This is useful with fruit, meats, other foods. It has been used to prevent freezer burn in fish and stickiin candy.

ectins in Alcoholic Fermentation. Alcoholic fermentation of fruits and is with yeast, usually Saccharomyces cerevisiae, yields ethanol, OH, and very small amounts of other organic compounds. Occaally methanol will contaminate the fermentation beer; and unless it is fully removed by distillation, it will be in the final product. This nanol arises by demethylation of pectins by pectin esterase.5 Yeast does form an enzyme capable of hydrolyzing pectins, and consequently the tion does not commonly occur in fruit fermentations. But pectin rase is abundant in the peel of citrus fruits and also in fungi. The acy of the enzyme rises as the pH increases from 1 to 6 and the producof methanol goes up. If citrus fruit is limed before fermentation, or if or grain with relatively high pH becomes contaminated with molds, amount of methanol formed may be fairly high.

ms and Mucilages

number of plants form gums on their bark or fruit when they are indor when they are stung by insects. An exudate forms which contains gh per cent of polysaccharide. This can be readily gathered and dried will absorb water and form a sticky mass. Many natural gums are commercially and used as adhesives or thickeners. Gum arabic from acacia tree, gum ghatti, gum karaya, and gum tragacanth are com-

eaweeds and a number of seeds contain polysaccharides which are lar in composition to those produced as exudates and which are used in lar fashion. Alginic acid and its salts are now widely used in the United es as thickeners. Agar is produced from seaweed (Gelidium algas) and igeenan from Irish moss (Chondrus crispus). Locust bean gum and

ice seed gum are examples of those from seeds.

or the most part, fruit gums, on hydrolysis, form pentoses, hexoses, are nic acids. Table 3.6 shows the distribution of monosaccharides and

nic acid in gums.

um Arabic. This gum is produced on the bark of Acacia trees when they win not, dry, and high locations. It is extensively used industrially as a kener and emulsifying agent and has, therefore, been the subject of a iber of studies. Gum arabic occurs as a calcium salt with an equivalent the between 1000 and 1200. The branched molecule appears to have a backbone of galactose residues held 1-3 with other monosaccharides uronic acids in the side chain.

where:

Galp = Galactopyranose
Araf = Arabofuranose
Rhap = Rhamnopyranose
GA = Glucuronic acid
(Rhamnose is 6-deoxy mannose)

Gum arabic is widely used, particularly in confections, where strong stab gels are needed.

Seaweed Polysaccharides. The seaweed polysaccharides are produced large amounts from various seaweeds throughout the world. Some se weeds have a relatively low cellulose content and a high per cent of oth polysaccharides. Often the surface of the plant is covered with these corpounds which give it a slippery feel. They are readily extracted by boiling water or alkali and appear on the market in either a very crude state or patially purified. In some parts of the world they are used for food, but in the United States they appear in foods and other products as thickeners are adhesives.

Although many of these polysaccharides have been used for centuries be man, most of them are not as yet well explored chemically and only tent tive or partial formulas can be assigned.

Agar. Agar, which is produced by boiling red seaweed in water, is linear galactan sulfate. The galactose units are held together with 1-3 link with about every tenth link 1-4. The sulfate occurs in the ratio of 1:3 galactose, as the calcium salt of the galactose ester. Both D- and L-galacto occur in the molecule. Agar has an enormous ability to absorb water are form gels at low concentrations. The Japanese have used it as a food for centuries although it is questionable if it has any food value for man. In the United States the food industry uses large quantities as a stabilizer are

TABLE 3.6. APPROXIMATE COMPOSITION OF SOME PLANT GUMS*
(Moles of Constituent per Repeating Unit)

Tum	D-Glucuronic Acid or Methoxyl Derivative	D-Galactose	D-Mannose	1 - Arabinose	D-Xilose	t-Rhamnose
Arabic	1	3	nil	2	nil	1
Damson	1	2	1	3	ca. 3%	nil
Cherry	1	2	1	6	ca. 3%	nil
Egg Plum	2	6	nil	7	1	nil
Purple Plum	1	3	nil	3(?)	1(?)	nil
Almond Tree	1	3	nil	4	2	nil
Lemon	+	+	nil	+	nil	nil
Orange	+	+	nil	+	nil	nil
Grapefruit	+	+	nil	+	nil	nil
Cholla	+	+	nil	+	nil	+
Mesquite	D-Galactu- ronic Acid	+	nil	+	nil	-

^{*}Trom Pigman, W. W. and Goepp, R. M., "Chemistry of the Carbohydrates," Academic Press, Inc., New York, N. Y., 1948.

gelling agent. It is used in icings on baked goods, in cream cheese, and in truit and vegetable jellies. Agar is often used in place of gelatin in jellied candies and marshmallows. It makes a fine jellied glaze for meat or fish.

Alginic Acid. Alginic acid or sodium alginate, is produced from the giant kelp of California, $Macrocystis\ pyrifera$, by extraction with sodium carbonate. It is a polysaccharide made up of units of mannuronic acid, joined β 1–4. It is used in ice creams, sherbets, syrups, processed cheese, salad dressings, and many other food products.

Carageenan. Carageenan is isolated from Irish moss by extraction with hot water. It is a mixture of polysaccharides made up of galactose monopor disulfates. The composition of a few of these polysaccharides has been studied and they appear to be almost linear with one unit branching. The crude carageenan is also widely used in food products. Most chocolate milk in the United States is stabilized with carageenan, and it is often used in milk desserts and other products.

Locust Bean Gum. This is added to ice cream powders, pudding powders, preserves and jams, salad dressings, and pie fillings, to increase the viscosity even gel the product. It is effective in low concentrations.

DENTIFICATION

Color Reaction

Carbohydrates can be identified in mixtures of biological materials by a number of color reactions. One of the most widely used general tests is the

Molisch reaction which is carried out by adding a drop of an alcoholic s lution of α -naphthol to a solution of the unknown. Concentrated sulful acid is then slowly poured down the side of the tube so that a layer formed below the unknown solution. In the presence of saccharides purple color develops at the interface. Carbohydrate acids or amines do n give the test, but mono- and oligosaccharides as well as many polysa charides are positive. The sulfuric acid reacts with carbohydrates to for furfural or one of its derivatives. These compounds produce colored proucts with phenols and phenol derivatives. Alpha-naphthol used in the Molisch test can be considered a phenol derivative.

Some of these color tests can more or less differentiate groups of carbohydrates from one another. Ketoses, pentoses, and uronic acids ca be differentiated from aldohexoses since they react more readily and often form different colors with phenols and their derivatives. Seliwanoff' reagent is prepared by dissolving resorcinol in alcohol and adding hydro chloric acid to not more than 12 per cent. Fructose, a ketose, will give red precipitate in 20 to 30 sec, while aldohexoses react much more slowly The formation of a red color with glucose after boiling for some time is be lieved to be caused by a transformation of glucose to fructose. The colo is the result of the formation of a furfural derivative and its subsequen reaction with resorcinol. Sucrose will give a positive test since it hydrolyze in the acid solution. Variations of the method include substitution of th 12 per cent hydrochloric acid by alcoholic sulfuric acid or acetic acid Phloroglucinol forms a cherry red color rapidly with pentoses, and slowly forms a dark red color which changes to brown with hexoses. Orcino forms greenish blue colors with pentoses and is somewhat more valuable than other reagents for differentiating pentoses and hexoses. All of thes tests must be used with caution and the time of heating and concentration

of reagents carefully controlled. In Tauber's test benzidine in glacial acetic acid produces a cherry red color with pentoses and glucuronic acid, while

Benzidine

Aniline Acetate

Naphthoresorcinol

hexoses give yellow to brown. Aniline acetate on filter paper turns bright red when held in the fumes from a mixture of pentoses and hydrochloric acid but not when hexose is present. The volatile compound is furfural. Uronic acids give a purple color when they are boiled with hydrochloric acid and then treated with naphthoresorcinol. These colored compounds are very soluble in ether and can be readily extracted from aqueous solutions. Anthrone is used in quantitative tests to give colors whose depth is proportional to the amount of carbohydrate present.

Anthrone

Sucrose can be identified by a color reaction when the solution is free of ratilinose, gentiobiose, and stachyose. It reacts with an alkaline solution of diazouracil to form a green color.

The ease with which mono- and disaccharides which contain free or potential aldehyde and ketone groups are oxidized gives methods for differentiating them from sucrose, which has no potential aldehyde or ketone groups, and from polysaccharides which have few reducing groups in a large molecule. Many reagents have been devised using copper, silver, mercury, and other metal ions which are reduced to the free metal or oxide by the aldehyde or ketone groups of sugars. Benedict's reagent, which is composed of copper sulfate, sodium citrate, and sodium carbonate, is widely used for the detection of glucose in biological fluids. The deep blue color of the copper ion complex changes to the yellow or to the brick red color of cuprous oxide, Cu₂O, precipitate when glucose is present. Other substances which are readily oxidized can interfere; but when Benedict's solution is used cautiously, it is a very valuable reagent.

Some colored organic compounds which on reduction change colors also used in detecting reducing sugars. Yellow picric acid changes to de

red picramic acid. Orthodinitrobenzene, methylene blue, and safranin a other reagents.

Dilute solutions of iodine are used to detect some polysaccharides; an although it is difficult to differentiate them by this reaction, the methodoes have some usefulness. Amyloses give a blue color while amylopecting give violet. Glycogens give red brown and dextrins form colors from viole through reddish to colorless depending on their molecular weight.

The ability of iodine in neutral, slightly acid, or slightly alkaline solutions to oxidize aldoses can be used to determine them quantitatively. If the conditions are carefully controlled, aldoses can be quantitatively determined in the presence of ketoses which do not react.

Reaction Products of Mono- and Disaccharides with Hydrazines

The products of the reaction of mono- and some disaccharides with hydrazines are useful in differentiating many of them. Phenylhydrazine is the most commonly used reagent although dinitrophenylhydrazine and others sometimes form products which are easily isolated and purified. One molecule of phenylhydrazine reacts with a molecule of the carbohydrate to form a phenylhydrazone and when the product is highly insoluble as in the case of mannose, it precipitates. If the hydrazone is soluble, further reaction occurs and the osazone is formed.

Fermentation with Yeast

Some groups of carbohydrates can be differentiated from others by the ability of yeast (Saccharomyces cerevisiae) to ferment them with the formation of carbon dioxide and ethyl alcohol. Ordinary bakers' yeast will ferment D-glucose, D-fructose, D-mannose, D-maltose, and sucrose, but forms little or no carbon dioxide if supplied with D-galactose, lactose, of the pentoses. If the yeast is grown on a medium rich in D-galactose, the yeast then develops the ability to ferment the galactose. However, with ordinary commercial yeast, it is very easy to carry out a fermentation test and differentiate these sugars. Other strains of yeasts and other micro

organisms have been used to differentiate sugars and even to measure them quantitatively in carbohydrate mixtures. For example, D-galactose can be determined by the method of Wise and Appling in carbohydrate mixtures through the use of Saccharomyces carlsbergensis, a yeast which ferments it, and Saccharomyces bayanus, a yeast which does not. A table on the fermentation characteristics of a number of microorganisms is given by Pigman and Geopp.²²

Separation by Chromatography

In many foods complex mixtures of carbohydrates exist and attempts to demonstrate the presence of one carbohydrate when others are also present is often difficult. Separation of the carbohydrates by chromatography allows the application of reactions for their identification without interference. When small samples are available, paper chromatography is usu-

ally used; but if the samples are large and if it is desirable to isolate the pure or relatively pure carbohydrates, column chromatography is used.

The ion exchange column can be used with ionized derivatives of carbo hydrates. Thus the products of reaction of carbohydrates with borates ar anions and are taken up by anion exchange resins. (See Boeseken.⁴)

Optical Activity

When solutions of pure mono- or disaccharides are available, the determination of the optical rotation serves to identify or to determine quantitatively the amount of a sugar present. This method is applied to ray and purified cane and beet sugars but is not very useful in natural food where complex mixtures occur. A polarimeter is used in measuring optical activity of a solution containing an accurately weighed sample. The specific rotation is calculated from the formula:

$$(\alpha)_{\rm D}^{20} = \frac{100 \ \alpha}{l \times c}$$

where $(\alpha)_D^{20}$ is the specific rotation at 20°C, and, using the D line of the sodium spectrum as a source of light, α is the observed rotation, l is the length of the polarimeter tube in decimeters, and c is the weight of the sugar in grams per 100 ml of solution. When the sugar is known, since its specific rotation is also known, the equation can be rearranged to solve for the concentration of the sugar:

$$c = \frac{100 \,\alpha}{l \times (\alpha)_{\rm D}^{20}}$$

The method can sometimes be applied to mixtures of carbohydrates where the rotation can be made to vary under uniform conditions. For example the Clerget method for sucrose in the presence of glucose and fructose depends on the fact that hydrolysis of sucrose results in a change in rotation. The rotation of the sample is measured before and after hydrolysis and since the specific rotations of sucrose and of the equimolar mixture of glucose and fructose formed on hydrolysis is known, the percentage of sucrose can be calculated.

$$S = \frac{100(P - P_1)}{133 - 0.5(t - 20)}$$

S is the percentage of sucrose while P is the original rotation of the mixture, P_1 is the rotation after hydrolysis and t is the temperature in degrees centigrade. Because fructose is unstable in acid solution, enzymic hydrolysis is sometimes used.

Other Quantitative Determinations

The most widely used chemical methods for the quantitative determination of sugars depend on their ability to reduce alkaline solutions of metals such as silver, mercury, bismuth, and copper. Of all the methods which have been devised, those depending on the reduction of cupric ions to cuprous oxide are the most widely used. A monosaccharide such as glucose reacts with mild oxidizing agents to form a large number of products. The aldehyde is oxidized to a carboxyl group, but the alcoholic hydroxyls are also capable of oxidation. If conditions are right, the chain may be completely fragmented and a number of moles of cupric ion reduced. Since this is true, it might appear that such a reaction would be an unlikely prospect for the development of a quantitative method. Nevertheless, when conditions of reactions and reagents are standardized, surprisingly accurate results can be obtained. The amount of cuprous oxide formed is proportional to a given amount of glucose under standardized conditions. The cuprous exide is determined by numerous methods. In some procedures it is weighed directly, in others it is ignited and changed to cupric oxide and then weighed. It can be dissolved and determined volumetrically by adding a ferric salt and titrating with permanganate, by adding a definite amount of iodine and titrating the excess iodine with thiosulfate, as well as with thiocyanate and silver salts. In some procedures the cuprous oxide is dissolved in the reaction mixture and determined directly. For example, in some colorimetric methods the cuprous oxide reduces molybdic acid to blue molybdic oxides. The depth of color is proportional to the amount of cuprous oxide formed and hence to the amount of carbohydrate present in the unknown. There is also a method which depends upon the determination of the excess amount of cupric ion remaining in the reaction mixture.

In some methods the sugar solution is titrated directly into the boiling copper reagent and the end point—when all of the cupric ion has reacted is determined by the disappearance of the blue color, by spot tests, or with methylene blue as an internal indicator.

Other methods depend on the oxidation of the sugar molecules with ferricyanide in the presence of a rather high pH. Here again the extent of the oxidation depends on the nature of the reagent, the alkalinity, the temperature, and the length of time used. But again, if the conditions are standardized, the reduction of the ferricyanide is proportional to the amount of carbohydrate present. Here, too, reducing substances other than the reducit g sugars interfere with the determinations and give high results. During the reaction, ferricyanide is reduced to ferrocyanide. The amount of ferricyanide remaining can be measured by reducing it with an iodide and measuring the amount of iodine released with standard thiocyanate. In some

procedures the sugar solution is directly titrated with the ferricyanide be use of an oxidation-reduction indicator such as methylene blue. Here again, conditions must be carefully standardized in order to calculate the concentration of the reducing sugar from the amount of ferricyanid reduced.

A number of color reactions of monosaccharides have been used for the quantitative estimation of these sugars. A sensitive micromethod which has been widely used is the color reaction with anthrone. The carbohydrate it treated with sulfuric acid to form a furfural derivative which gives a colo with anthrone. The depth of the color is proportional to the amount of carbohydrate present under standard conditions. Either a colorimeter or a spectrophotometer is used to measure the color.

Carbazole and some of the phenols such as orcinol (3,5-dihydroxy-toluene) which are used in the qualitative detection of carbohydrate have been used in the development of quantitative methods.

Iodine in dilute alkaline solutions will oxidize aldoses without markedly affecting either ketoses or nonreducing sugars and this reaction can be used in determining quantities. Many other compounds will reduce iodine and the method must be used with care and under strictly standardized conditions. But when compounds are separated in chromatograms, the method can be used for the quantitative determination of aldoses. Iodine in alkaline solution forms hypoiodate which reacts with the aldose. Excess iodine is then back titrated by acidifying and titrating with standard thiosulfate.

CHANGES OF CARBOHYDRATES ON COOKING

Carbohydrates in food show some general changes on cooking or processing of the food. In general, these may be summarized as:

(1) Solubility

In some cookery processes soluble carbohydrates are dissolved. This physical change is usually of importance in mixtures where sucrose is one of the components.

(2) Hydrolysis

Some polysaccharides are hydrolyzed during cooking or processing. This action is particularly noticeable in foods which contain enough acid so that they are sour to taste and consequently are called acid foods. Pectic substances undergo this hydrolysis with the result that the fruit or vegetable becomes mushy and the juice thickens as the pectic and pectinic acids disperse in it.

Hydrolysis of starches also occurs to a limited extent during cooking. Occasionally the extent of hydrolysis may be so great that the thickening power of the starch is decreased. Thus when lemon or cherry pie filling is nickened, the mixture must not be cooked too long after the starch is added or the viscosity decreases again. Dextrinization of starch occurs on he crust of bread during baking but this is not simple hydrolysis. The starch molecule is degraded and the reactions which occur are much more complicated than hydrolysis.

3) Gelatinization of Starch

When starch is mixed with cold water, no apparent change occurs; but vhen the water is heated, the viscosity of the mixture increases and if the concentration of the starch is sufficiently great, a gel is formed. Often if the tarch suspension does not gel at the elevated temperature, it will when the nixture is cooled. This process is called the gelatinization of starch. It depends on a number of factors. The temperature at which gelatinization starts and the exact changes during the course of gelatinization are characteristic of, first, the variety of starch. Thus wheat and cornstarch show different behavior patterns, and even floury cornstarch from the crown egion of most common corn acts differently from horny cornstarch from he region of the endosperm. (See p. 76). Second, the pH at which gelatiniration is measured is most important. This fact was not recognized during he early work on starches and consequently the determinations of gelatiniration made at that time are sometimes worthless. Third, the temperature it which observations are made and the length of heating are important. And fourth, gelatinization is influenced by the size of the granule. The temperature of gelatinization decreases as granules decrease in size.

The microscopic appearance of the starch granules changes markedly on neating, in three stages. The first stage, in cold water, is marked by the mbibition of approximately 25 to 30 per cent of water. This is apparently reversible effect because the starch may be dried again with no observable thange in structure. The viscosity of the starch-water mixture does not hange during this phase. The second stage occurs at approximately 65° C or most starches, when the granules begin to swell rapidly and take up a arre amount of water. Thus Meyer and Bernfeld¹⁹ report that cornstarch kes up 300 per cent water at 60° C, 1000 per cent at 70° C, and at the oint of maximum swelling 2500 per cent, based on the original weight of he starch. The granules change in appearance during this second phase, and some of the more soluble starch molecules are leached out of the granule. The liquid surrounding the granules will show a color with iodine even hough the granules are still intact. This stage is not reversible. The third

stage is marked by more swelling. The granule becomes enormous, often void is formed, much more starch is leached out, and finally the grant ruptures, spilling more starch out into the surrounding fluid. The viscos of the fluid increases markedly, and the starch granules stick together that they can no longer be picked apart.

The swelling of starch, particularly amylose, which results in an increasin viscosity of a starch-water mixture and the formation, under proper conditions, of a gel is now believed to occur through the binding of water. In starch granule, amylose and amylopectin molecules are loosely bound to gether by hydrogen bonds of the hydroxyls. A hydrogen on a hydroxyl one molecule is attracted by the negative charge of the oxygen of a hydroxyl on another molecule, and this attraction forms a weak link between the molecules, as shown in Diagram A.

These aggregates of molecules, held together weakly, form micelles. As the temperature of a water-starch mixture rises, hydrogen bonding decreases for both the starch-starch bonds and water-water bonds and the size of the particles diminishes. The tiny water molecules begin to freely penetrate between starch molecules. Conversely, as the temperature decreases, water molecules are bound between the starch molecules (Diagram B) and there is an increase in size or swelling. Where two starch molecules were originally bound together, there are now the two starch molecules with water molecules in between.

The sticking together of granules is believed to be the result of molecule from adjacent granules becoming attracted and enmeshed in one another.

Gel formation occurs through the formation of a three-dimensional network of starch molecules, particularly the long straight chain amylos molecules. (See p. 126.) These molecules become interlaced through attractive forces between the molecules and particularly through hydrogen bonding on water molecules. Highly branched glycogen does not form gels occurstals, and gel formation in starch is consequently believed to be primarily the function of amylose rather than amylopectin.

Starch gels are readily cut by shearing forces and reduced to liquid. The are consequently called *thixotropic* gels. This phenomenon is often important in food preparation since stirring disrupts the gel. On standing the ge forms again.

On aging, most starch gels show marked syneresis or weeping in which he water gradually passes out of the interstices of the gel.

Another process also occurs both in gels and in viscous but still liquid ispersions. Part of the starch aggregates and forms microcrystals that prepitate. This is called the retrogradation of starch. Thus most starch dispersons which have stood for several days or weeks contain a deposit at the attom of the vessel. The process also occurs in gels and is believed to be ne of the important factors in the staling of bread. Figure 3.8 (see page 4) attempts to show the change in organization of molecules which occurs a starch precipitates.

RUDE FIBER

Mangold¹⁶ defines crude fiber as the sum of all those organic components f the plant cell membrane and supporting structures which in chemical nalysis of plant foodstuffs remain after removal of crude protein, crude at, and nitrogen-free extractives. Thus the crude fiber should be composed f the cellulose, hemicellulose and some of the materials that encrust the ell walls such as lignins and pectic substances. Actually the quantity of rude fiber found is a function of the method used in its determination. Trude fat is removed by ether extraction while crude protein is dissolved by hydrolyzing with dilute acid. During this hydrolysis, other compounds will likewise be hydrolyzed. However, when a standard method is used, remoducible results can be obtained and a given food will fall within a range of values. Crude fiber values for a given food show the same variation with limate, soil conditions, and degree of maturity as do other values.

The method which has been adopted by the Association of Official Agriultural Chemists employs boiling the sample in 1.25 per cent sulfuric acid, o lowed by boiling in 1.25 per cent sodium hydroxide. The residue is composed of the crude fiber and some salts. The crude fiber is determined by trying to constant weight, weighing, and then ashing, and weighing again.

The difference between the two weights is the crude fiber.

Interest in crude fiber determinations centers either in the detection of dulteration or from the nutritional standpoint in the bulkiness of the diet. ome foods are occasionally adulterated with inert material by unscrupus sellers. For example, spices may be adulterated and consequently exaded with finely ground sawdust or with the waste from the spices. Crude per determinations will detect this increase. Mustard has a range of crude but by the official method of 4.2 to 6.5 per cent, while cloves ranges from to 9 per cent. If the crude fiber determination is much out of line, the roduct has probably been adulterated.

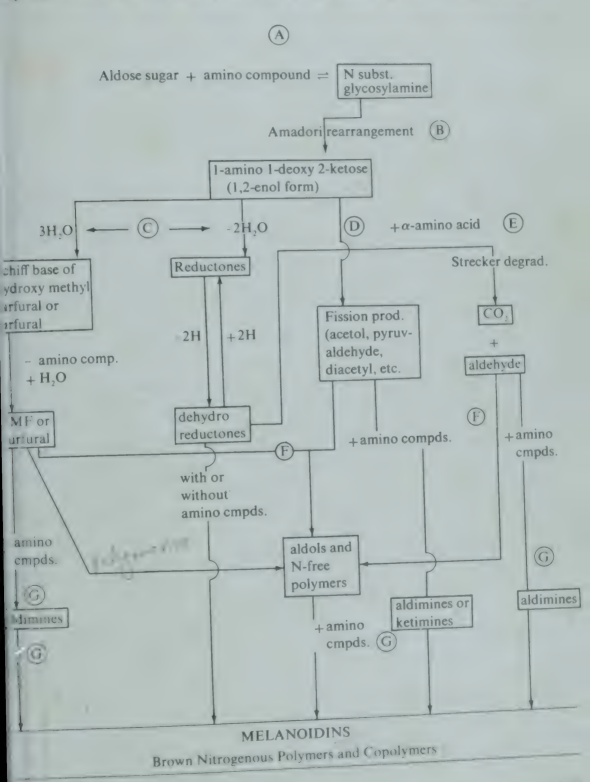
The compounds in crude fiber make up most of the bulk or residue of diet, and are not hydrolyzed by the digestive fluids of human beings. Hever, bacteria present in the paunch of ruminants and even those present the colon of human beings may bring about considerable hydrolysis. fibers that escape hydrolysis are excreted in the feces. They serve a very function here, however, because most of them are capable of absorb water, and they render the feces soft enough to pass out of the bereadily and bulky enough to induce defecation.

BROWNING REACTIONS

The browning reactions are complex reactions which occur when ma foods are processed. In some the brown flavor is highly desirable and intimately associated in our minds with a delicious, high-grade produ In coffee, maple syrup, the brown crust of bread and all baked goo potato chips, roasted nuts, and many other processed foods controll browning is necessary. Yet in other foods, browning during processi is undesirable and forms off-flavors and dulled or even objectional colors. In drying fruits or vegetables and in canning or concentration orange juice, it is highly desirable to avoid browning. The presence carbohydrates in foods is intimately connected with the browning which occurs. Other compounds are sometimes important, but they are on which have some of the reactive groups of the reducing sugars and which are similar to them in their chemical properties. The pigments which a formed are high molecular weight polymers whose constitution is difficu to determine. The browning reactions appear to be complicated not only to the final product but also as to the course of the numerous reactions. has been exceedingly difficult to assess the chemistry of this change in the complex mixtures encountered in almost every food. During the past fif years, and particularly during the last twenty, study of the browning rea tion has been carried forward by the use of model systems. In this type study one, two, or sometimes three compounds are allowed to react and the intermediates, products, and course of the reaction followed. Even th method of excluding and simplifying has not yielded all of the answers b any means, since the possible reactions are numerous. To a student n completely familiar with the field, the many investigations appear at first be wholly unrelated and the state of the problem all confusion. However, 1953 Hodges¹² attempted to correlate and integrate knowledge which ha accumulated up to that time about browning reactions. An attempt will I made to briefly review his ideas.

In the past, three general types of browning reactions have been reconized to occur in foods during processing: (1) The reaction of aldehyd

nd ketones, among them the reducing sugars, with amino compounds uch as amino acids, peptides, and proteins. This is independent of the resence of oxygen. (2) Caramelization, the change which occurs in polydroxycarbonyl compounds such as reducing sugars and sugar acids when nev are heated to high temperatures and which is also independent of



oxygen. (3) The oxidative change of polyphenols to di- or polycarbo compounds and possibly the oxidation of ascorbic acid. This may be tially or wholly enzymatic. No matter what the type of reactants the bropigments, called melanins or melanoidins, are unsaturated polymers. It will notice that in each of the broad types of reaction except the third carbonyl or polycarbonyl compound is important, and in the third the formation of carbonyl compounds. Since the actants of the third type furnish carbonyl compounds, it is correct to consider these compounds indispensible to any type of browning reaction. When a food is extracted to remove carbonyl compounds, browning is tarded or eliminated. It is therefore believed that although the course these three types of reactions are incompletely understood, they are reactions of importance in the browning of foods.

Hodges has given a scheme for browning reactions and has included three general types in one scheme. See the diagram on page 109. Browing may occur by compounds entering the scheme at any point. Althouthe first reaction, is a reaction of an aldose sugar with an amino co pound, there is some evidence that ketoses can cause browning and they may react in a similar fashion. This first reaction of an aldose wan amine is shown in the following Amadori reaction:

$$\begin{array}{c} RNH \\ HCO \\ (CHOH)_n + RNH_2 \longrightarrow (CHOH)_n \\ CHOH \\ CH_2OH \\ Aldose \\ Amine \end{array}$$

$$\begin{bmatrix}
RNH \\
HC \\
(CHOH)_{n+1}
\end{bmatrix}
+
RNH \\
HC \\
CH_2$$

$$COH$$

$$CH_2$$

$$CHOH)_n$$

$$CH_2OH$$

The product of the Amadori rearrangement can then undergo a numb of fates depending on the conditions of the reaction. It can, in neutral acid media, lose water and form a ring compound of the Schiff's base of h droxy methyl furfural, or furfural, and then eliminate the amine to for

rm reductiones. These are compounds with a high reduction potential at have not been identified as yet. Or finally, the product can undergo sion to form small molecules such as acetol, CH, COCH, OH; pyruvaldede, CH, COCHO; diacetyl, CH, COCOCH,; and others. D' In the

heme shown (page 109). Hodges has indicated that all of these carbonyl ompounds react with amines to form aldimines or ketimines. G or polynerize to aldols and similar large molecules which subsequently react with mines. F The final brown pigments are products that contain nitrogen. his scheme covers all of the reactions of the first broad general type of rowning reactions—the reaction of aldehydes and ketones with amino ompounds.

The second broad general type of browning reaction is caramelization. his type has not been as extensively studied. It is, of course, a reaction hich does not require an amine. Sugars will show caramelization when eated to relatively high temperatures. This type of reaction is markedly fected by high pH, as is the first general type. While the browning of nese carbohydrates is not as rapid as in the presence of amino compounds, is accelerated by the presence of carboxylic acids, the salts of these acids, hosphates, and metallic ions. These accelerators are commonly present in ods. The nitrogen-free intermediates formed in carbonyl-amino browning eactions are also produced in nonamino browning. The formation of 1,2 polization, furfural and hydroxymethyl furfural by dehydration, and sugar ssion products has been demonstrated in a few model systems. It has also een shown that these intermediates will form colored polymers in the bsence of amino compounds. So in general this type of browning reaction ts into Hodge's scheme, joining it after the Amadori rearrangement and orming different (nitrogen-free) polymers.

The third type of browning reaction is also believed by Hodges to have, the least, a possibility of fitting into the scheme. This is the browning much occurs in the presence of oxygen when fruits or vegetable tissues are ut or bruised. It is completely discussed in Chapter 7 on the fruits and egetables. Enzymes bring about the transformation of tyrosine, catechols, rother polyphenols to quinones. Since catechols are ortho di-phenols, are therefore conjugated enediols and are reductones. Hodges con-

siders that they enter the scheme at this point. Not a great amount known of the reactions of quinones as they brown. Browning may be

matic, but some steps may be nonenzymatic and may well pass through actions common to the other intermediates in types (1) and (2).

Browning is a problem of general interest in food technology. The lanumber of studies now being conducted both on model systems and food products makes it possible that these reactions will soon be maken completely understood.

We see then that both carbohydrates and amino compounds are imptant in nonenzymatic browning. Habib and Brown⁸ have tested the valid of this hypothesis in potato chips which are notable for their variation browning. Browning during frying is desirable in potato chips not only the effect on appearance but also for the flavor which the browning retion imparts. Habib and Brown⁸ separated the amino acids and sugpresent in four varieties of potatoes by paper chromatography. The patatoes were studied at three different times and their browning reaction compared to the content of carbohydrates and amines: 20 days after havest, after 4 weeks of cold storage at 40° F, and after a further reconditioning for 4 weeks at 75° F. The reconditioned potatoes had the lowest level amino acids, with the drop in quantity particularly in the basic amino acids. These potatoes also produced the lightest-colored chips, supports the hypothesis that browning depends on reactions of carbohydrates amino acids.

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CHAPTER FOUR

Proteins in Foods

WHILE PLANTS ARE ABLE TO UTILIZE inorganic sources of nitrogen stammonia, nitrates, and nitrites, man and other higher animals are formost part dependent on a source of amino acids to build their body teins. Although some bacteria can utilize atmospheric nitrogen, the ganisms are low in the scale of life and far removed from man. I animals are directly or indirectly dependent on plant protein. Often plant protein is consumed by one animal, digested and synthesized its proteins, and reaches man after passage through one or more and Thus, large fish rely on plant foods by way of a long line of smaller smaller marine animals down to the tiny ones actually eating the food.

Every living cell contains protein. Indeed, the name "protein" of from the Greek root which means "to be first." In man synthesis of many proteins that make up body tissue requires the presence of a acids. Approximately 20 different amino acids are in the body proteins although certain proteins such as thyroglobulin contain one or mortusual amino acids as well. All 20 of the amino acids are not require the diet since, as has been demonstrated, man is capable of synthemall of the amino acids except 8 (leucine, isolaucine, lysine, valine, three tryptophan, phenylalanine, and methionine).

$$\begin{array}{c} \text{CH}_3 \\ \text{CH}_2 \\ \text{CH}_3 \\ \text{CH}_2 \\ \text{NH}_2 \\ \text{Leucine} \end{array} \qquad \begin{array}{c} \text{H}_2\text{CH}_2\text{CH}_2\text{CH}_2\text{CHCOOH} \\ \text{NH}_2 \\ \text{NH}_2 \\ \text{Lysine} \end{array}$$

$$\begin{array}{c} \text{CH}_3 \\ \text{C}_2\text{H}_5 \\ \text{NH}_2 \\ \text{NH}_2 \\ \text{Isoleucine} \end{array} \qquad \begin{array}{c} \text{CH}_3 \\ \text{CH}_3 \\ \text{CH}_3 \\ \text{CHCHCOOH} \\ \text{NH}_2 \\ \text{Valine} \end{array}$$

wever, some source of nitrogen in the amino form must be available, amino group from other amino acids enters the nitrogen pool and can sed for the synthesis of any amino acid whose carbon skeleton can be e. Man's food must supply sufficient amino acids for the purpose of ding proteins. It must allow for occasional synthesis of nonessential ro acids, as well as supplying adequate amounts of the essential amino s that he cannot synthesize.

TEINS IN MAN'S DIET

Ithough man's need for protein and its amino acids is rather stringent, method of fulfilling his need varies considerably in different parts of world. Primarily the difference has depended on the availability of is. In the tropics many peoples have developed dietary patterns based varily on plant foods with the cereals most abundant. In most of these animal protein in the form of meat, fish, eggs, and milk is added navailable. Although it is difficult to build an optimum diet on plant lucts alone, it is possible if the diet is varied.

any of the proteins present in plant tissues are deficient in one or c of the essential amino acids. For example, zein, one of the proteins orn, is lacking in lysine and tryptophan; while gliadin, one of the its of wheat, is low in lysine. However, both wheat and corn contain proteins that possess these amino acids. A diet restricted entirely to at or to corn is low in lysine and, under the stringent demands for d ynthesis, is inadequate. If the wheat or corn is supplemented with ems that are relatively rich in lysine, then the amino acids supplied adequate.

Although the food mixtures eaten by primitive man and handed do diet patterns were probably adequate nutritionally, modern man doe appear to have an instinct for fulfilling his nutritive needs. Where he stricted to a vegetarian diet, malnutrition is often very com Kwashiorkor, the deficiency disease of many children in Africa and Latin American countries, is a protein deficiency complicated with deficiencies. In the part of Africa where this disease is common, yam the staple food of the people, while in Latin America it is corn.

In the Arctic regions the diet of the Eskimos is principally animal. only are fish and meat the principal food, but almost the complete ar is eaten, often uncooked. This diet is unusually high in protein and assortment of amino acids in these proteins meets the requirements.

the synthesis of protein in man.

In the temperate zones most diet patterns include both animal vegetable products. The common pattern when the economic level allowadequate in the amount of protein as well as in the distribution of amino acids. Inadequacy is almost always the result of economic faculthough it is occasionally caused by faulty eating habits.

CHEMICAL AND PHYSICAL PROPERTIES

Proteins are high molecular weight compounds that yield amino acide their principal hydrolysis product. The structure of proteins and the classification of proteins on the basis of solubility and structure is quately described in textbooks in organic or biochemistry and can be viewed there. A few of the properties of proteins which are not always studied in organic chemistry and a brief discussion of the methods of termining molecular weight and testing homogeneity will be presented.

Proteins are large molecules with some reactive groups, such as carband amino, on their surfaces. Because of the size of the molecules, they not form true solutions in water. When solubility of a protein is not ioned, dispersibility is intended. A dispersion of a protein in water a sequently has the properties of a colloidal dispersion rather than a solution. The reactive groups and the size cause a marked sensitivity only to the pH of the solution but also to the presence of many difference electrolytes. Protein molecules not only associate with small molecules also with one another, sometimes to form tightly bound and sometimes to some the size of the molecule is such that adsorption of proteins is extremal. Protein molecules are quite sensitive to many reagents and difficult. Protein molecules are quite sensitive to many reagents and ditions and readily undergo small transformations in structure. The

y of isolating a protein from its natural source, which occurs in a cell iological fluid, without any alteration in the molecule is great.

he field of protein chemistry is consequently a difficult one. Chemists learning how to handle better, determine, and evaluate large macro cules. Protein chemistry is profiting from work with other large moles and is also contributing to it. Despite the difficulty of working with eins, many investigators are actively engaged in the field and the num-of publications which appear each year, is very large. Organic chemists studying structure and reactions, physical chemists are attempting to e measurements and define the molecule, biochemists are describing ole of proteins in all aspects of life.

ne shape of protein molecules varies greatly. Some exist as fibers, others oheres while intermediate are many shaped like spindles, cigars, etc. manner in which a polypeptide chain can achieve a spherical shape or itermediate one, is the object of considerable speculation at present, iously the chain must be folded or coiled in some fashion and there to be some bonds which hold the chain in a relatively permanent shape, insiderable amount of study is at present directed to the solving of this olem.

ecular Weights and Homogeneity

olecular weights of proteins are so large that the classical methods for determining the molecular weights of compounds that will form solutions or vaporize are useless. New methods have been devised and relding significant information. The most widely used are those of electhoresis and sedimentation, with some work on osmotic pressure and a on light scattering. These methods also give some information on the ogeneity of a protein preparation.

production of a crystalline protein is no more proof of the purity repreparation than is the preparation of any crystalline compound. How molecular weight organic compounds, determining the melting tof the compound is usually possible, and if the melting point is sharp not depressed by a mixed melting point, we are reasonably satisfied of arrity. Proteins do not melt. If they are heated, they undergo decomnon. Some other methods must therefore be applied in order to astrate the purity of a preparation.

able free amino and carboxyl groups, proteins are amphoteric. In the of alkali, they act as acids and achieve a negative charge through orization and neutralization of free carboxyl groups. But in the presolation, the amino groups react with hydrogen ions and assume posi-

pH and positively charged at low pH. It will also be remembered the some intermediate pH, called the *isoelectric point*, the total charge of protein molecule is close to zero, with the available amino and car groups neutralizing one another and forming zwitterions. In an el field the protein molecules in a solution at high pH will move to the tive pole (anode) since they are negatively charged, but at a low pH will migrate to the negative pole (cathode).

They will not move if the solution is buffered to the isoelectric point of proteins. Migration in an electric field at a definite pH is called electric phoresis and is a valuable tool in determining molecular weights of protein as well as in separating mixtures or proving the uniformity of a crystall or purified protein. The rate of migration in a constant electric field pends on the charge on the molecule, the size of the molecule and, certain extent, on the shape of the molecule.

Tiselius has devised an apparatus in which measurements can be made constant temperature and with a minimum of boundary diffusion buffered protein solution is covered by a layer of buffer solution; the work cell is immersed in a constant temperature bath, and the electrodes serted. Protein molecules that are alike will move at the same rate will form a sharp boundary in the cell. A colored protein such as he globin can be seen moving through the cell. However, most proteins are colored and other methods must be used to follow their path. Usually refractive index is used and in a Tiselius apparatus provision is made record the changes in refractive index photographically. In a mixture proteins such as that present in plasma, a series of peaks will be present in the record indicating the presence of a series of proteins moving different rates. When the method is used to demonstrate that only one lecular species is present in a fraction, it is necessary to determine

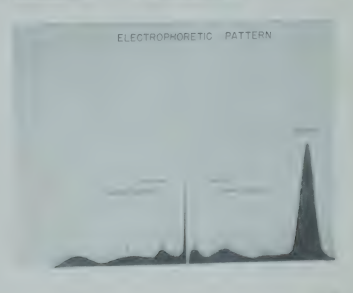
GURE 4.1. TISELIUS ELECTRO-LESIS APPARATUS. The elecvessels are on each side with higration chamber in the midconstructed so that fractions be cut out and migration fized. Courtesy of The Upjohn vanv.



HE TISELIUS ELECTROPHORE-PPARATUS. A light source through the sample and the tion of light gives a measthe protein passing. A cammounted at the left hand end tube and a power source is it. Courtesy of The Upjohn any.



TORE 4.3. AN ELECTROTIC PATTERN OF NORMAL I SHOWING THE PROTEIN TONS PRESENT. The ordinate e and the abscissa from the diffraction gives the amount tein. This is an oversimplized properties. Courtesy of The n Company.



electrophoretic mobility at several pH's. Occasionally two prot migrate at the same rate at a given pH when the *balance* of forces c the charge on one species of molecule, the weight of this molecule shape are exactly matched by that of the other species. This balance be impossible at other pH's and consequently, if the sample still sharp boundary at a second pH, it is considered pure.

Paper electrophoresis is widely used to demonstrate the presonumerous proteins in small samples of biological materials and also evidence of the homogeneity or heterogeneity of a given protein. A paper is wet with buffer solution and arranged so that one end is in of the buffer solution with the cathode and the other end is in a vess buffer solution with the anode. The protein is placed near one end strip. Under the influence of electric current, the protein moves paper; and if it is homogeneous, a thin zone will be maintained molecule moves at the same rate. However, a mixture of proteins see into zones. When the paper is suitably developed with reagents the color with proteins, the number of zones shows the number of posmetimes a sample must be run at more than one pH on separate paper to demonstrate all of the different proteins present.

Sedimentation. The ultracentrifuge has proved to be an important ment in determining the molecular weight of large molecules such teins. It is an apparatus invented in 1925 by the Swedish chemis Svedberg, for applying a very strong centrifugal force to a dispersive recent modifications of Svedberg's ultracentrifuge, speeds have be tained in which the rotor revolves at the rate of 60,000 times per with centrifugal forces developed equal to 500,000 times gravity (2 g). The rotor is operated in a high vacuum to minimize air resistancut down the amount of heat developed by air friction.

The rate at which a particle will travel in the field depends on the size, and density of the particle; on the density and viscosity of the persion medium; and on the centrifugal force. If the particles are identity will fall through the solution at the same rate and produce a boundary between the colloidal dispersion and the dispersion med mixture of molecules will form several boundaries depending on the ber of molecular species present. The method is consequently us proving the homogeneity or heterogeneity of a protein fraction.

A single boundary is evidence of the presence of molecules all same size. However, it is not certain that the protein is pure if it single boundary unless other methods support this. Occasionally pare found to give a single boundary in the ultracentrifuge but to mixtures under other conditions. The boundary is followed in the

aner that is used for electrophoresis, an optical system which records change in refractive index photographically. The apparatus which has devised to record this change is by no means simple.

olecular weights can be calculated from measurements of the variable edimentation. While estimates of the molecular weights by this means pare favorably with those of other methods, they are only approxime. When good agreement by several methods is obtained, confidence in results is enhanced.

fact has been used to determine the molecular weights of a number of eins. If a protein dispersion is separated from water by a semipermemembrane making osmosis possible, the water tends to migrate in ter amounts into the protein dispersion. However, osmotic pressure is of the properties of solutions that depends on the number of moles present. Thus since the number of protein particles is low, the ostic pressure is low. When the weight of the protein dispersed is known, then possible to calculate the molecular weight.

smotic pressure measurements of protein dispersions are technically er difficult. They are usually made at the isoelectric point of the pro-

The dispersions often attain equilibrium very slowly so that days must llowed for a single experiment. In this length of time it is difficult to ent denaturation of the protein and the growth of microorganisms. Strements by this method compare very favorably with other molecu-

e ght determinations and are of about the same accuracy.

ight Scattering. When a beam of light is passed through a colloidal ersion, part of the light is transmitted and part is scattered. This is well-known Tyndall effect which is seen with every airport beacon on y nights. The amount of scattering depends on the number of colloidal icles and also on their size. Hence a measurement of the amount of scattering gives a method for calculating the molecular weights of cir. particles. It has been used effectively in many protein fractions gives good agreement with other methods of determining molecular hts of large molecules. Its accuracy is of approximately the same r as other methods.

laivsis of Unusual Component. Occasionally a protein contains an an component which can readily be detected and estimated, and can serve to give a minimal molecular weight. For example, the of hemoglobin has been used for such a purpose. When other methowed values for the molecular weight approximately four times that I on iron determinations, it was apparent that each hemoglobin molecuntains four iron atoms. Since iron can be determined with great

accuracy, the molecular weights approximated by other methods hence be refined. However there are few proteins that contain an u element such as iron.

Chemical Properties of Proteins

Amphoterism of Proteins. The ability of proteins to react either as or as bases has been mentioned. The free carboxyl groups of protein very weakly ionized but, nevertheless, are available for reaction bases. The free amino groups, on the other hand, are hydrogen accand are available for reaction with acids. The number of these groups able for reaction on each protein molecule is small since it is on side chain carboxyls of glutamic acid and aspartic acid and the side amino group of lysine which can react. The carboxyl and amino group an amino acid such as glycine are tied up in the peptide bonds. The groups of histidine, tryptophan, proline, and hydroxyproline are rebasic as amino groups, but can accept hydrogen ions. The ability of protein to act as base or acid will depend not only on the number these groups, but probably on their placement, whether they are clothesurface of the molecule, or covered.

Binding of Ions. Proteins can bind both cations and anions through action with either the carboxyl group or the free amino groups. A values above the isoelectric point a protein exists as negative ion binds or reacts with cations. At pH's below the isoelectric point i exist as a positive ion and will react with or bind anions. The precedition of protein by heavy metal ions depends on the formation of these tein salts. A number of metallic ions have the ability to form coor tion compounds (complex ions) and these will readily form through tion with amino N, the substituted amine of the peptide link, and imid N. Copper, nickel, and iron are examples of metals which form these plexes.

In most foods mixtures of proteins occur, and at a given pH some teins may be below their isoelectric point and others above theirs. Uthese conditions cations will be bound by some of the proteins and a by others.

Hydration of Proteins. Proteins can form hydrates with water and type of reaction is often important in food chemistry. A protein modeontains a number of groups in which a nitrogen or an oxygen atom tains a pair of unshared electrons and is therefore capable of form hydrogen bond. The nitrogen in the peptide link as well as the nitrogen the free amino group are in this condition, and by their relative ativity can attract the hydrogen of a molecule of water.

double bonded oxygen of the carboxyl (_COOH) group or of the ronyl (CO) of the peptide link is more strongly negative and has a ter attraction for the hydrogen than the nitrogen. The ionized groups ned at pH levels below or above the isoelectric point have a greater afy for water and consequently hydrate formation is greater at pH's other the isoelectric point. The water molecule which has been bound can act another molecule of water since it possesses an oxygen with an unred pair of electrons. Aggregates of water can therefore build up around h polar group on the protein molecule. In protein dispersions in which er compounds that form hydrates are present a condition frequently urring in food chemistry -- competition for water may occur between the tecules. Electrolytes, sugars, alcohols, and many other substances have tendency to combine with water and form hydrates and may compete n protein for the water. The extent of hydration of a protein dispersion efore depends not only on the concentration of the protein dispersed also on the pH, the presence of other substances which combine with er, and the temperature.

'recipitation with Antibodies. Although it is now possible to determine approximate molecular weights of proteins, there is no chemical hod for differentiating most protein molecules. If a protein contains an sual element such as the iron present in hemoglobin, it is possible to that to differentiate the protein from those which do not possess iron. ny proteins are very similar in molecular weight and amino acid comition but are nevertheless different structurally. So far, the only method precise identification of proteins is a biological reaction. If a foreign tem (antigen) is injected into the blood stream of an animal, a sub-(antibody) which precipitates that protein is developed. For example, rabbit is given several injections of egg albumin, an antibody will deon in the blood of the rabbit and later, even a minute amount of egg mi will result in the formation of a precipitate. These antigen-antibody mons are highly specific. The blood of the rabbit which is highly sensiegg albumin will not give a precipitate with any other albumin. This st mean that egg albumin differs structurally from all other albumins n though their molecular weights and chemical and physical properties similar. Thus, although no chemical methods have been developed as

yet to demonstrate subtle differences between protein molecules, biologic tests are able to establish them.

Native and Denatured Proteins

The proteins that occur in the tissues, whether within the cells or in the fluids, of living plants and animals are called *native* proteins. These larges molecules are quite fragile and many reagents and conditions cause slight of extensive changes in the structure of the protein. The properties of the protein change and this, of course, indicates an alteration in the structure of the molecule. The resultant changed protein is called *denatured protein*, and it usually shows a very different solubility from the native protein since the change in the structure allows the molecules to aggregate and precipitate Frequently the denaturation is irreversible, but occasionally under very mild conditions reversible denaturation occurs.

Denaturing agents include acids, alkalis, alkaloids, heavy metal salts and other compounds such as urea and ethanol. Most of these are not important in food chemistry although the possibility of denaturation of protein must never be overlooked. Many ions such as iodide, bromide, and chloride are denaturation agents and the synthetic detergents are active at low concentration.

Although denaturation in foods is usually the result of increase in temperature, sometimes denaturation can be produced by mechanical means. Extensive whipping of an egg white foam will produce some denaturation and the foam will begin to break as the protein starts to precipitate. Ultraviolet light, high pressures, and ultrasonic vibrations are all denaturation agents. Spreading of protein films at interfaces usually results in denaturation.

In studies of protein structure where the native protein is the center of interest, it is imperative that denaturation be avoided. This is often exceedingly difficult during the tedious separation of protein from other substances in the tissue.

Reversible denaturation of protein occurs under very mild conditions or very short exposure and may occur in living systems. Reversibly denatured protein is able to return to the native state and show the original physical properties and biological activity. If the protein is rapidly removed from the reagent or condition that causes denaturation and placed in a solution at the isoelectric point where its charge is at a minimum, rapid precipitation occurs. But if it is placed in a solution with a pH different from the isoelectric point, it will develop a charge, molecules will repel one another, and it will rapidly return to the native state.

The nature of the transformation that occurs when a protein is denatured

now believed to be an unfolding of the molecule. It may well be that in versible denaturation only a limited unfolding occurs while in irreversible enaturation it is more extensive. It scarcely seems probable that a long olypeptide chain which has become completely disentangled would readily able to reassume the spatial relations of the native protein. Many studs have attempted to find differences other than solubility between native and denatured protein. The sulfhydryl (SH) and disulfide (SS) oups are readily detected in proteins and have been extensively studied. ften the denatured protein shows a different content of sulfhydryl groups and some years ago it was thought that the process of denaturation was indamentally one in which sulfhydryl groups are formed from disulfide. oday the results are more commonly interpreted to mean that more ilfhydryl groups are exposed as the molecule unfolds. There is also some vidence that more of the phenolic groups present in tyrosine, and more the indole groups in tryptophan are available for detection in deatured protein. These observations are also interpreted as exposure on unolding of the molecule.

In many respects native and denatured protein are very similar. But in ome of the biological properties which are more discriminating of structre than the methods of chemistry now available, marked changes are observed. In general denatured proteins are more readily attacked by roteolytic enzymes. When the protein studied is an enzyme or an antiody, denaturation is accompanied by a loss of activity. This surely indi-

ates a change in the molecule.

Since details of protein structure are still not completely known, it is not arranged arranged arranged and the protein is a oiled or folded molecule in which certain parts of the polypeptide chain t together with definite other parts, it is not unlikely that denaturation expresents some disorganization of this pattern.

sel Formation

Gel formation is a very important process in food chemistry. Not only o the properties of living cells, both animal and vegetable, depend on the el structure, but in food preparation the stiffening which occurs during neat and flour cookery, the rigidity of pectin and starch gels, the high iscosity of many plant juices, the changes that occur in egg cookery and carry other processing operations are a function of the gel.

Although colloid chemists have studied gel formation for many years, and although the literature is full of papers describing gels, the number of compounds investigated is relatively small compared with the number bout which we should like information. As in many other topics in food

chemistry, we must thank the colloid chemist for the work and insignar gained and then attempt to apply the limited results to a large ber of situations, where not only has the gelling compound escaped but the number of other compounds present produces a very compl situation. Therefore, it is highly worthwhile to review briefly the p state of knowledge concerning gel formation and the applications to chemistry now known.

A gel is a remarkable phenomenon, displaying the property of rig sometimes at quite low concentrations of solute and yet often showing properties of the solvent practically unchanged. For example, most in water show vapor pressure and electrical conductivity very clowater. Gel formation occurs in gelatin dispersions in water with as a concentration as 1 per cent gelatin and in plasma with a fibrinogen centration of 0.04 per cent. The phenomena displayed by gels are contant and are not always the same with different gels.

When colloidal dispersions of some relatively large molecules are co the viscosity increases to a point at which some rigidity is attained. point is called the *gel point*. Usually on further cooling or sometime standing, the rigidity of the gel increases. Many gels on continued stan lose solvent and the gel shrinks in a process called *syneresis* or *wee* Gels differ considerably in their rigidity. Some are deformed under pres and others even flow.

Theories of Gel Formation. Many attempts to explain the properties gel have been made, and through the years a number of theories have proposed and argued back and forth. Three major theories are now ported by colloid chemists. With some gels one theory is more likely another. These theories are: (1) adsorption of solvent, (2) three-dimensinetwork formation, (3) particle orientation.

- (1) Adsorption of Solvent. This theory postulates that adsorption of vent molecules by the solute particles results, on cooling, in the format of larger and larger particles with increasing layers of solute. The larged particles eventually touch or overlap enclosing more solvent, so the entire system is immobilized and rigidity occurs. Support of this the depends on demonstrating that adsorption of solvent molecules is very tensive and that the adsorption increases with decreased temperature.
- (2) Three-dimensional Network. This theory postulates that the compound capable of gelation is either fibrous in structure or can react itself to form a fiber. On cooling the fibers form a three-dimensional work by reacting either at widely separated intervals on the chain of relatively small distances. The bonds established which tie the fibers the three-dimensional network can be either primary bonds between f

mal groups, secondary bonds, such as hydrogen bonds, or nonlocalized condary attractive forces such as might occur between alkyl groups. The first type of bond would yield a network that would possess conerable permanence. This type could only occur in gels that are not readily sociated by cutting or beating, for example. They would be capable of elling and shrinking, within limits, because it would be possible for the ers to be pushed farther apart without disrupting the network, by aightening the chain of atoms. There would, of course, be a limit to the nount of solvent that could be taken up in swelling, or the amount that uld be lost. (b) A secondary type bond is one of considerably lower ength than a primary valence bond, and these gels would consequently broken by relatively small forces. A gel that can be disrupted by beate might possibly have a network structure where secondary valence ands hold the fibers together. (c) The third type of bond, a nonspecific traction between portions of the molecules or along the entire molecule, n explain gelation in a few situations. The properties of this type of gel ald depend on a nice balance of forces the solute molecules would atact one another at some spots but would be separated by solvent moleles that are attracted at others. The result would be gelation. If the solnt molecules were too strongly attracted to the solute, then the solute olecules would have no opportunity to contact one another and a netork would be impossible. Under these conditions a gel would not form. n the other hand, if the attraction of the solute molecules for one other were too strong, dispersion in the solvent would not occur. A ec pitate or an insoluble compound will be formed rather than a gel. his theory can explain gel formation in systems that are very demandg as far as temperature, concentration, pH, and salt concentration are ncerned.

Particle Orientation. This theory postulates that in some systems are is a tendency for the solute and solvent particles to orient themves in definite spacial configurations through the influence of long range rees such as occur in crystals. Certain protein crystals have the ability take on or lose water without distortion of the crystal. They may form uctures of this type. Tobacco mosaic virus has been studied and found form gels under a large range of concentrations. X-ray diffraction studies die ate that the gels possess a two-dimensional lattice. This is interpreted near that the particles are oriented in one direction.

The gelation of a few proteins has received considerable study in the sc. but many have as yet not been investigated. Gelatin and fibrin clots we probably received the largest share of attention, while myosin and omyosin from muscle, and denatured egg albumin have also been stud-

ied. Ferry⁷ finds that the three-dimensional network theory of gel tion most closely fits the data that have accumulated for proteins.

Gelatin. Gelatin is a partially degraded protein. It is prepared from collagen of skin, ligaments or bone by alkali or dilute acid hydrowhen bone is used, the bone salts are usually dissolved with acid the protein then hydrolyzed with Ca(OH)₂, lime. The native protein are hydrolyzed are collagens (see p. 176), and it is usually assumed the collagen molecules are broken into shorter fragments that are new less still fibrous in structure. A study of gelatin from bone incompared that the collagen is first hydrolyzed into molecules of approximation molecular weight and then more slowly hydrolyzed at random smaller fragments. The gelatin is therefore composed of an assortment molecular species and cannot be considered a single compound. The sortment of molecules apparently occurs in all gelatin samples althoug assortment will not be identical in each sample.

Gelatin gels have been studied extensively for many years and much have been accumulated concerning the properties of these gels under conditions, some applicable to food chemistry and some not. The fa which influence the rigidity of a gelatin gel include concentration o gelatin, temperature, molecular weight, added reagents, and pH in ranges—although the influence of these factors is limited. The rigid roughly proportional to the square of the concentration. The proporti ity is not strictly valid for some samples and under all conditions. perature has a marked effect and the rigidity increases rapidly with creasing temperature. This relationship is linear. Molecular weight l direct effect on rigidity so that samples that have high average mole weights form more rigid gels under controlled conditions than those lower average molecular weights. The effect of pH is only slight in range close to the isoelectric point of the sample. Thus a series of ge 3 per cent concentration showed little change in rigidity over a pH r of 4.6 to 8.2, while at a concentration of 1.5 per cent, constance rigidity occurred at pH 4.3 to 6.7. The influence of salts on rigidity been studied but in ranges of concentration of both gelatin and salt w are of little significance in food work. For example, a series of isoele gelatin gels with a concentration of 14 per cent showed a decrease in r ity with increasing sodium fluoride concentrations up to 0.5 M and the creasing rigidity.

Denatured Protein Gels. The gels of denatured protein that have studied show very marked effect by many conditions. They form under very specific conditions. There have been only a limited number definitive studies of denatured protein gels, and those on denatured

Bumin are perhaps most numerous. In all of these gels the dependence on H and salt concentration as well as on the presence of nonpolar solvents idicates that the gel is produced by a type 3 (see p. 127) three-dimensional twork. The nice balance of forces that is necessary to achieve gelation dicates that the molecules must show a relatively nonspecific type of ataction between the polypeptide chains. There must be a balance between rees of attraction in part of the chain and forces of repulsion in others, he solvent must neither be too strongly nor too weakly attracted to the nain.

The conditions under which egg albumin has been studied have not been rectly applicable to food preparation. One comprehensive study used propyl alcohol and water as the solvent, another used relatively high conntrations of acetic acid in water. But all studies indicate the sensitivity falbumin gels to various conditions of pH and salt concentration. There spears to be two steps in the formation of the gel (1) denaturation and 1) gelation. In the denaturation reaction a fibrous molecule is formed. It is followed by the setting up of a network through the attraction of portions of molecules to one another.

ETERMINATION OF PROTEIN IN FOODS

Accurate determinations of the protein content of the complex mixtures high are encountered in foods is a difficult job. Kirk¹² in his discussion the "unsatisfactory state of analytical development in this field" makes e following statement: "The reasons are apparent. Proteins form a very verse group of similar compounds of extraordinary complexity, with idely different compositions and properties, yet difficult to separate cometely, to purify and to dry. Their amphoteric nature, high adsorptive caucity, hydration properties, and sensitivity to electrolytes cause them to try widely in behavior depending on the composition, pH, and temperare of the solvent medium."

In the determination of proteins in food, interest is commonly on the tal protein content of the food, rather than on an accurate determination the presence of a specific protein. Occasionally in some research probnit it is of interest to determine specific proteins. An example is the agen content of various types of meat, or the total amount of gluten able of development in various grains. Often rough determination of tal protein content is sufficient for the study. This is well and good, if a rough protein determination is not then used as if it were an accurate termination.

Since proteins are so readily precipitated from solution, it might concluded that it would be relatively simple to precipitate the protein f a food, filter it off, dry, and weigh it. But the problem is far from sim It has not been possible so far to find a satisfactory reagent that precipitate all proteins free of contamination with other substances. ides, salts, and many organic molecules are carried down in protein cipitates. Some can be removed by washing, but the process is difficul achieve quantitatively and is laborious and time consuming. Drying protein to constant weight is at present impossible. Proteins have a h degree of hydration and the sample slowly loses water over a long per of time. Although the hydration is a reversible reaction, complete remo of water is very difficult, if not impossible, without protein decompositi The freeze-dry (lyophile) method has so far not been widely applied to p teins but may be a possible method. So far, direct methods of estimat protein show considerable variation and present difficulties in the rep duction of consistent data.

A few methods have been investigated for precipitating protein and the measuring the amount of precipitating agent that has combined with a protein under the conditions of the experiment. None of these methods free from a considerable margin of error; and none has been wide adopted.

One method of determining protein is analysis for carbon. After all of ganic compounds except protein have been removed as well as possible analysis for carbon can be attempted. With a few biological systems it is been successful. Analysis of sulfur has not been used extensively. The most widely used method of determining protein is the analysis in itrogen.

When a food is analyzed, provision must be made to remove nitrogen containing compounds other than proteins. A cell or biological fluid contains many such compounds. Free amino acids, some of the lipids, uncreatine and creatinine, thiamine, heme, and many other compounds contain nitrogen. While all of them are present in relatively small amount it must nevertheless be remembered that they as well as protein contributed to the amount of nitrogen present in a food, and that a change in the concentration of one of these components will alter the amount of nitrogen present as surely as a change in protein.

The number of grams of protein in a food is often calculated multiplying the number of grams of nitrogen by 6.25. This constant is or rived from the assumption that proteins contain 16 per cent nitrogen a 100/16 = 6.25. The assumption is not valid since all proteins do not contain exactly 16 per cent nitrogen. Protein is often reported as "cru

compounds which are nonprotein and the "protein constant."

eldahl Method

The usual method employed for the determination of nitrogen in foods the *Kjeldahl method* and various modifications have been devised to prove its accuracy and speed. It is an oxidation of organic compounds by furic acid to form carbon dioxide and water and the release of the nitron as ammonia. The ammonia exists in the sulfuric acid solution as ammium sulfate, but the carbon dioxide and water are driven off. Sulfur oxide is the reduction product of the sulfuric acid and it too is volatile.

Organic compounds + H₂SO₄ → CO₂ + H₂O + (NH₄)₂SO₄ + SO₂

ne digestion of the sample to form ammonium sulfate is the most diffilt part of the operation. Although considerable effort has been expended obtain accurate analyses, many factors cause incomplete formation of amonia or loss of some of it from the digestion mixture. Attempts to minate or minimize these errors have centered around studies of: (1) emical form of some of the nitrogen, (2) addition of salts to digest mixtee. (3) use of oxidizing agents, (4) catalysts, (5) length of time of distion, (6) use of reducing or hydrolyzing agents prior to digestion, (7) ntamination with ammonium salts used in fractionation. Kirk concludes at the most important sources of error are connected with the catalyst, time of heating, and the addition of reducing and oxidizing agents.

Many catalysts have been used to speed up the decomposition of the mple. Copper, mercury, and selenium are most widely used. Mercury apars to give better recovery of nitrogen than copper, but it must be preditated before the ammonia is estimated since it forms a complex with ammonia. Selenium shortens the clearing time of the sample, but it has en given a false sense of completion of digestion when clearing time and mpletion have been confused. Many workers have used 1.5 times the aring time as the length needed for complete digestion, but it has been monstrated that even this length of time is not always sufficient. Chibll, Rees, Williams, and Jonnard have advised 8 to 16 hours of digestion or clearing in order to insure completion. With a long digestion time, a quantity of acid present must be carefully watched. If the amount falls low, decomposition of NH4HSO4 occurs with the loss of ammonia the mixture.

ombinations of catalysts have been used effectively. Copper, mercury, d selenium are used together as well as copper and selenium, and mercy and selenium.

Numerous oxidizing agents have been added at the end of the diperiod to complete the oxidation. Potassium permanganate was the reagent used and in recent years persulfates, perchlorates, and hyperoxide have been used. On various types of samples losses of nithrough the formation of amines or of free nitrogen have occasionall reported with all of these reagents except hydrogen peroxide. Howe has not been tested with a wide number of different types of professional provides must be used cautiously since it is commonly prewith acetanilide to depress the formation of oxygen and water, an nitrogen in the preservative will act as a contaminant of the detertion.

The chemical form of the nitrogen is important. Thus the terminal a group on lysine is difficult to release as is the imidazole nitrog histidine and tryptophan. Sufficient length of time must be allowe complete transformation into ammonia. At the same time the amou acid must be controlled so that it does not fall sufficiently to allow loss of ammonia from the mixture.

Lysine Histidine Tryptoph

Salts such as sodium or potassium sulfate are commonly added to digestion mixture in order to raise the boiling point of the mixture consequently shorten the time of digestion. However, the ratio of sa acid must not rise too high or the loss of ammonia by decomposition on NH₄HSO₄ will occur. Some workers use phosphates in place of the fates, with good results. Here, too, the ratio must not be too high.

Measurement of the ammonia after it is formed by digestion is calculated out by a number of different methods, all of them quite accurate. In the ammonia is distilled off after the addition of large quantities of and trapped in a known quantity of standard acid. The acid is then titrated to determine how much ammonia has distilled.

Another method uses the direct estimation of the ammonia by rea with Nessler's reagent to form orange Nessler's salt.

$$2K_2HgI_4 + NH_3 + KOH \rightarrow NH_2Hg_2I_3 + 5KI + H_2O$$

The Nessler's salt is insoluble and precipitates; but if the amount is s it can be produced in colloidal form and estimated colorimetrically

me degree of accuracy. Technically, the method is often difficult to carry it because of the tendency of the colloidal dispersion of Nessler's salt to come cloudy and then precipitate.

A method which has not been as widely used is oxidation of the amnia with hypobromite and back titration of the hypobromite with line.

$$2NH_3 + 3NaBrO \rightarrow N_2 + 3H_2O + 3NaBr$$

umas Method

The Dumas method is a combustion method for the estimation of nitron in a sample, which has seldom been used on food samples. Comparins of Kjeldahl methods with Dumas have occasionally been made. Often e Kjeldahl method gives nitrogen values a little lower than the Dumas. the Dumas method the sample is heated in a combustion train in the esence of carbon dioxide. The gases are passed over hot copper gauze so at any nitrogen present as an oxide is reduced to free nitrogen. Carbon saide and water are absorbed and the nitrogen gas formed measured rectly.

When a food sample is analyzed for nitrogen, the accuracy of the method analysis. Kjeldahl or Dumas, varies with a number of factors, but often e error exceeds 1 or 2 per cent and sometimes even 5 per cent. Since the ta are then multiplied by a factor that is approximate, the final answer nnot be exact and gives only the crude protein. Often the nitrogenntaining compounds other than protein are neither removed nor estimated, and, therefore, cannot be subtracted from the results. Most of our ta have then this further approximation. Data on the protein amount of ods must, therefore, always be taken as approximate only, unless a reful check is made to assure that all approximation has been eliminated d that the protein constant actually applies.

mino Acids

Amino acids are often determined in proteins. Although most proteins, diparticularly those of animal origin, contain all 20 of the common bir o acids, the amount in the protein is frequently of some interest. Ice the essentiality of some amino acids have been recognized, a great more of studies have been made of the level of feeding one amino acid the requirement for another. Although these data are in the province utrition, there is also a considerable interest in amino acid distribution the field of food chemistry.

The problem of separating compounds as closely related chemically as amino acids and estimating the concentrations of each is by no means all. In recent years the development of the technique of chromatography

and the rapid increase in knowledge of microbiological assays have tated the solution of this problem.

Chromatography was studied in 1861 by Schoenbein and used is separation of chlorophyll from other plant pigments by Tswett in The term "chromatography" is applied because Tswett separated compounds, mainly on a calcium carbonate column. When the meth applied to compounds which possess no color, it is still called "chrotography."

The technique depends on the differential adsorption of compo even those which are closely related chemically, by a solid adsorbent. mixture of compounds can be dissolved in a suitable solvent and p through a column of the adsorbent. As the solution trickles through compounds may move and be adsorbed at differential rates. Thus one pound which is strongly adsorbed will be held at the top of the col while one which is poorly adsorbed will be at the bottom. Sometime of the compounds are strongly held at the top of the column, but they be separated by the use of a second solvent which detaches them erentially. When the second solvent is poured over the column, the fraction collected at the bottom contains the substance held least stro and most readily dissolved in the second solvent, while subsequent f tions will contain other members of the mixture. The difficult part of technique is finding those solvents which will effect separation, as wel a suitable adsorbent for the column. This is empirical and can be very t consuming.

Paper chromatography is an application of this technique to a filter pa strip and has frequently been applied to the separation of amino acids mixture of compounds is placed at one end of a strip of filter paper. paper is suspended in a cylinder and dips into an organic solvent sa rated with water. The atmosphere in the vessel is likewise saturated v the vapors of the two solvents. The filter paper has a stronger affinity water than for the organic solvent and the components of the mixture m up the strip at different speeds. When the solvent front reaches the top the paper, the paper is removed, dried, and sprayed with some compound that will react with the components to form colored products that can seen. The components will be found at different places on the strip. T is ascending chromatography, but descending chromatograms can made by dipping the top of the strip in a trough of solvent-solvent m ture and allowing the components to move down the strip. Twomensional separation can be achieved by running the mixture in one dir tion, then turning the paper 90° and running again. Better separation sometimes possible by this device.

Chromatography can be used to separate even quite small samples of ano acids after hydrolysis of the parent protein. The quantities of ano acids present then can be estimated by various methods.

In acid hydrolysis of any protein, tryptophan is destroyed and humin ormed. Humin is a black or dark brown precipitate which always acapanies acid hydrolysis. In the presence of carbohydrate, it is especially indant. Since most foods contain carbohydrate along with the protein, nin formation presents an error in quantitative estimations of amino d content. The most satisfactory method is to remove carbohydrate, is is often very difficult to achieve without loss or change of the protein. *Quantitative estimation of some individual amino acids* can be carried out specific reactions to give colored compounds or precipitates. The

specific reactions to give colored compounds or precipitates. The kaguchi test, for example, is a test for arginine which can be applied ter to proteins or to amino acid mixtures. It is carried out by treating sample with α-naphthol and sodium hypochlorite. In the presence of tinine a deep red color develops. The method may be used quantitatively ce the depth of the color is a function of the amount of arginine present example of a precipitating reagent is Reinecke salt—a complex to f chromium, NH₄[(NH₃)₂Cr(CNS)₄] ammonium diamminetetratinatothiohromium III—which forms a precipitate with proline and droxyproline. Other reagents are discussed in textbooks of biochemistry

in special books on proteins and their analyses.

The microbiological methods are fairly recent developments which have n very valuable in analyzing mixtures of amino acids because of the ec and reproducibility of results obtained. The methods depend on preing a nutrient medium which contains all of the compounds essential the growth of a particular microorganism except one amino acid, which I be assayed. Addition of this amino acid results in growth of the croorganism in proportion to the amount of the amino acid added. Cule tubes are set up and graded amounts of the unknown are added to a ies of tubes. Standards are set up at the same time with graded amounts he pure amino acid. The unknown can be compared to the standards by asuring the rate of growth of the microorganism. Sometimes turbidity he medium after a 24 hr period of growth in the incubator is used. th organisms which form acid such as some of the lactobacilli, titration he acid formed can be used as a measure of the number of cells present. on'y difficulty in carrying out these determinations is in finding an .nism which requires a particular amino acid. Streptococcus faecalis reres nine amino acids and can be used in the estimation of these. Pure tures must be used and all of the techniques necessary for handling .roorganisms without contamination observed.

HEAT TREATMENT

For many years it has been recognized that the nutritive value teins changes with heat treatment, and a great many studies have devoted to the problem. Unfortunately the "nutritive value" of a prostill rather poorly defined because it is complicated by many for "Nutritive value" is usually measured by comparing the growth resin a species such as the albino rat to graded amounts of a purified por some food source. The growth response depends on numerous besides the level of protein consumed: (1) the amino acids present protein or proteins, (2) the quantity of each essential amino acid post (3) the previous nutrional history of the test animal, (4) the extent gestion of the protein, and (5) the rate of digestion of protein. Resit has been shown that other factors such as mineral and vitamin let the diet, the presence of rancid fat, starvation, and hormones such as sone all influence the rate of growth and the response to specific leverotein.

It is well established that heat treatment can alter the nutritive value a protein when measured under constant conditions in a number of sp. Many proteins show a decrease in value, but in a few there is an incommodate proteins, meat proteins, blood globulin, cereal proteins, fish coconut meal are among the numerous products which have been structured in some studies the purified proteins are used and in others the food.

The chemical changes that take place on heating have been investion both isolated protein systems and in foods and have been found to sist of three types of reactions: (1) the reaction of protein with car drate, resulting in the destruction of some amino acids and a change digestibility of the protein by proteolytic enzymes; (2) reaction of p in the absence of carbohydrate which results in a decrease in the avaity of an amino acid; and (3) heat inactivation of enzyme inhibitors as ovonucoid.

When proteins are heated with carbohydrates, particularly the mon disaccharides, interaction of the protein with the carbohydrate occur continued heating, browning and destruction of amino acids occurs, simple system where only amino acids and glucose are used, there is a gressive loss of histidine, arginine, valine, and leucine in the order as shown by microbiological assay. The same type of reaction has bee served in numerous protein-carbohydrates systems as well as in foods. For example, evaporated milk has a lower nutritive value measured by rat growth, than fresh, whole milk. And if the evaporatik has been subjected to heating before feeding, it suffers a further

onstrated that the reaction occurs through the amino groups of the tem since if these groups are acetylated and rendered unavailable for non, the protein loses its power to bind carbohydrates.

ot only has it been shown in some studies that amino acids are deved, but the protein also becomes resistant to hydrolysis by some teases. For example, heating casein or wheat gluten with glucose causes cance to hydrolysis of the protein by trypsin or papain and to a less ked degree by pepsin, chymotrypsin, or pancreatin (which contains a ture of proteases).

ieating protein in water or alone also affects the amount of amino acids the resistance to enzymatic hydrolysis of the protein. Denaturation is to make a protein more susceptible to enzymatic hydrolysis. In one is was shown that if casein or ethanol-precipitated lactalbumin is ted, hydrolysis by enzymes is not as complete. Soluble lactalbumin did show this behavior.

namy foods carbohydrate is closely associated with protein and on heattnere is a reaction between these compounds. In one study it was wn that on heating, sunflower meal, peanut meal, cotton seed meal, corn are decreased in biological value, while beef (no carbohydrate lent) is unchanged and linseed meal is improved. However, in another by canning or heating beef decreased its biological value. Canning scauses a small decrease in the amount of methionine and a slight rease in resistance to digestion.

reprovement in nutritive value occurs when enzyme inhibitors are deved. A number of enzyme inhibitors have now been found in foods. On ucoid, one of the proteins of egg white, was the first recognized. This is the activity of trypsin. Although it is rather resistant to heat the temperatures used for most cooking and food-processing operations, as be destroyed by sufficient heat. It is significant that this antistic factor is apparently ineffective in the human, although it appears to retracted from soybeans is found to depress growth of rats on diets that the protein source is either casein or soybean protein. This into destroyed by heating in steam.

beans and their reaction to various types of treatment have been early studied. (See Almquist!, Swanson and Clark!, and Rice and I he data of Melnick and Oser! indicate that methionine is libely very slowly from heated soybeans although digestibility is not

affected. They suggest that the low biological value may result the tardiness of methionine to enter the metabolic pool while amino acids are available in good supply. Other investigators who measured the digestibility of heat-treated proteins as well as their b cal value for growth promotion, favor the view that the molecul sorbed from the gut are not free amino acids but derivatives, unavfor protein synthesis.

The significance of these observations in human food is questio Low biological value can only be demonstrated in animals on low p diets. Since the diets eaten by man usually contain proteins from sources, these proteins can supplement one another. Toasting bread decrease the amount of some of the amino acids in the wheat pr but when it is supplemented by egg or milk at the same meal, the loss is insignificant. When men eat restricted diets during times of fa or in areas of poverty, the small loss could have significance.

PURE PROTEINS FROM SOME FOODS

Plant Proteins

The study of the proteins of plants has been plagued through the by the same difficulties that the study of any protein entails. There is difficulty of isolation of a protein from the complex mixture which all occurs in cells, without modifying the molecules. Many proteins occills conjugated with lipid, carbohydrate, and other molecules, but those conjugated proteins which are not readily dissociated are isolated. With plant proteins there is the usual trouble encountered in dling a large, relatively reactive molecule such as a protein. Most protein are readily denatured by a wide variety of reagents and conditions, and utmost care must be used to avoid reaction. 3, 13

In view of the tremendous number of plants in the world, and the view of the tremendous number of plants in the world, and the view in cells with structure, the amount of knowledge concerning proteins that has accumulated is relatively meager. See Table 4.1. spermatophytes (flowering plants) have received the greatest amount attention while the gymnosperms (conifers, etc.) have been scarcely sturble seeds commonly used for food have been investigated most frequency see Table 4.2. The thallophytes, which include algae, fungi, actinomy and bacteria, have had considerable study. This has occurred becauthe interest of the brewing and baking industries in yeast and becauthe rapid development of the field of antibiotics, which are produce some of these plants. Much of the old work is incomplete and of questions.

A BRYOPHYTE, AN ALGA, FUNGI, AN ACTINOMYCETE, BACTERIA, AND PHYTOPATHOGENIC VIRUSES* TABLE 4.1. AMINO ACID COMPOSITIONS" OF BULK PROTEINS IN SPERMATOPHYTES, PTERIDOPHYTES,

					I hallophytes	phytes		Phyto-
Amme Acid	Spermato- phytes ^h	Prerido- phytes ^b	Bryo- phyte ^b	Algab	Fungi	Actino- mycete	Вастепа	pathogeme
Glycine	0.4						1	1.4- 2.1
Alanine	4.4- 5.1	-	1	1	-		1	4.5- 6.1
Valine .	3.3- 4.5				2.2- 4.0	4.6	3.5- 5.1	4.5- 6.6
Leucine }	7 , 7.3				2.7- 4.9	4.5	3.4- 5.1	5.8- 6.0
Isoleucine	7.1 3.6	1	1	1	1.1- 4.0	1.9	3.3- 4.7	3.0- 4.3
Phenylalanine	2.4- 2.6	1	-	1	1.1- 1.9	1.5	1.4- 2.2	2.8- 5.1
Cyst(e)ine	1.1- 1.6	1.1-1.2	1.3	2.2	1	ı	1.5- 1.6	0- 0.5
Methionine	1.2- 1.6	1.5-1.7	1.4	1.2	0.4-0.9	9.0	0.6- 1.5	0.0- 1.2
Tyrosine	2.3- 2.7	2.1-2.6	2.4	1.3	1.1- 2.3	1	0.8- 2.2	1.7- 3.1
Tryptophan	1.4- 1.9	1.1-1.4	1.7	1.0	0.6- 1.0	0.1	0.3- 0.6	0.4- 1.8
Arginine	12.4-14.0	15.3	1	ı	4.9-13.3	10.3	4.8-12.2	17.5-20.5
Histidine	3.6- 4.0	1	1		1.4- 7.4	2.5	1.2- 4.4	0.0- 1.2
Lysine	5.0- 6.8	6.4	*	1	3.9- 9.1	4.6	5.7-10.6	1.6- 2.9
Aspartic acid	4.7- 5.4	1		1		1	1	8.0-8.8
Glutamic acid	6.4- 7.8	1		1	4.9- 5.8	1	4.0- 7.3	3.7-8.9
Proline	3.1	1	1		-	1		4.0- 4.3
Hydroxy proline					1			c· +
Serine		1	[1	1			4.6- 7.5
Threonine	3.0- 4.0	1	1	1	2.0- 3.7	3.1	2.4- 3.6	4.5- 6.7
Amide	4.7- 6.0	4.9-5.3	5.5	5.6	1	1		0.0

As per cent protein nitrogen.

"Main photosynthesizing tissues

*From Lugg. J. H., "Plant Proteins," Ichances in Protein Chem. 5, 230-304 (1949)

able validity, in the light of recent developments but, nevertheless, of s interest.

Work so far seems to indicate that the bulk proteins of plants, ticularly among the spermatophytes, are very similar in amino acid c position. Seeds, however, appear to possess proteins which are quite similar in composition. A few thallophytes do not fall in line with analyses of tissue proteins of the rest of the plant world, but these data r to be reinvestigated.

Milk Proteins14

Milk is one of the excellent sources of protein in man's diet. Its role a source of protein has been emphasized frequently both in the scientific erature and in that for the general public. Its place as one of the Basic Se in the recommended diet plan for Americans stems from its fine contrition of protein as well as calcium and other nutrients to the diet.

The proteins of cow's milk have been extensively studied, and those human milk have received considerable attention. Very little is known the proteins present in the milk of mammals other than that of the cow human. In this book when the term "milk" is used, it refers to cow's milk

In recent years with refinements in the techniques of separating prote and in the development of physical measurements, electrophoresis, se mentation, osmotic pressure, and other methods, it has been possible to tend earlier work and with more confidence describe the proteins of mit These proteins are separated by precipitation at definite pH, salt fraction tion, and occasionally heat coagulation. These protein fractions must further purified in order to gain homogeneity.

Casein. Casein is the name assigned to the fraction precipitated by acifying milk to a pH of 4.7. It is present in cow's milk to the extent of to 3.5 per cent, in human milk 0.3 to 0.6 per cent. It may be further pufied by redissolving and precipitating again. Since it is very readily of tained, it has been studied for many years. The casein produced by the method was shown many years ago to be a mixture. Many attempts have been made to prepare pure proteins from this mixture, and it appears present to contain at least three proteins. These have been named α -, and γ -casein and differ from one another in their molecular weight their rate of migration in an electric field, and their phosphorus context. The amounts of some of the amino acids which have been determined lied wise show that these are different proteins. See Table 4.3.

Casein is also precipitated from milk by the action of the enzyme rem (the extract is called rennet) isolated from calves' stomachs; and the pro-

TABLE 4.2. APPROXIMATE PERCENTAGE OF AMINO ACIDS IN PLANT PROTEINS: PROTEINS OF SEEDS CALCULATED TO 16.0 g OF NITROGEN*

				and a or MALKOGEM.		
ro Acids	Cotton Seed Meal	Linseed Meal	Peanut Flour	Sovbean Meal	Oats	Rice
nine	7.4	6.9	9.9	5.8	_	_
atidine	2.6	1.9	2.1	2.3	6.0	7.2
sine	2.7	2.0			2.0	1.5
			3.0	5.8	3.3	3.2
rosine	3.2	5.1	4.4	4.1	4.5	5.6
yptophan	1.3	1.6	1.0	1.6	1.3	1.3
enylalanine	6.8	5.8	5.4	5.7	6.9	
stine	2.0	1.9	1.6	0.6 ± 1.4		6.3
thionine	1.6	2.3	1.3	2.0	1.8	1.4
reonine	3.0	4.5	1.5	4.0	2.3	3.4
ucine	5.0				3.5	3.9
		7.5 ± 2.8	5.5	6.6	8.0	9.0
leucine	3.4	3.4 ± 0.3	3.4	4.7	5.3	5.1
line	3.7	5.8 ± 1.3	4.0	4.2	6.5	6.4
ycine	5.3	_	5.6			_

Adapted from Block, R. J. and Bolling, D., "Amino Acid Composition of Proteins and Foods: Anacal Methods and Results," 2nd ed., Charles C. Thomas, Springfield, Ill., 1951.

m of discovering the difference between acid precipitated casein and min casein has been studied extensively. Conventional methods do not ow any difference in casein precipitated by either acid or rennin. They differ in the fact that rennin casein cannot be clotted a second time. The may indicate a difference in the structure of some fraction of the min casein. It has been found that the electrophoretic pattern of renninated casein differs from that of the original casein in that the α -casein ows two peaks, indicating the presence of two types of molecules. Some restigators believe that rennin hastens the dissociation of α -casein into a components. Since rennin is incapable of clotting milk if the calcium are first precipitated, in the past it has been customary to consider a reaction with acid or with rennin as following quite different pathways.

America it has been customary to call the native protein *casein* and the juble protein formed from rennin action, *paracasein*. The course of the action has been often written:

plincation. At present it is not possible to describe the course of the action.

Berridge' says that any theory of rennin action must account for the fact at there is a considerable time lag at low temperatures between the comet on of the action of rennin and the clotting. It must also account for

TABLE 4.3. COMPOSITION OF CASEIN FRACTIONS*

Constituent	α-Casein (Per Cent)	β-Casein (Per Cent)	Proba Alcohol Casein (F
Phosphorus	0.99	0.66	0.
Total N	15.5	15.4	15.
Amino N	0.99	0.73	-
Arginine	4.3	3.4	2.
Histidine	2.9	. 3.1	2.
Lysine	8.9	6.6	4.
Tyrosine	8.1	3.2	2.
Tryptophan	1.6	0.6	_

Mobility in Veronal buffer at pH 7.78 and ionic strength of 0.1 6.98 3.27 (cm²volt⁻¹sec⁻¹ \times 10⁻⁵

the fact that the concentration of calcium ions has little effect at centrations below 0.011 M and there is virtually complete precipitatio the casein at all concentrations above 0.0125 M. Neither the theory that action of rennin splits α -casein into two types of molecules, nor the reaction adequately explains the facts so far observed.

Whey Proteins. Milk contains approximately 0.6 to 0.7 per cent prowhich is not precipitated on acidification to pH 4.7. This represents at 20 per cent of the protein contained in skim milk. These whey protein were separated long ago into two fractions which were called globuling lactglobuling and albumin or lactalbumin. The old names still persist; but the light of modern technique and the demonstration that whey contain number of proteins, not just two, it is desirable to modify them.

Classical globulin is separated from whey by saturation with magnes sulfate. This fraction contains a number of compounds as shown by trophoresis. It has been possible to separate two homogeneous fract which are called euglobulin and pseudoglobulin.

The fraction which is soluble in saturated magnesium sulfate is classical albumin. A crystalline component which appears to be homogous has been prepared from this fraction. It has been named β -lactobulin since it appears to be identical with the β -fraction from uncentrifuge studies of milk serum.

Other fractions have been purified and more than two peaks occur in electrophoretic pattern of whey. There is, therefore, the probability these components occur in milk. They are still to be fully characterized.

Colostrum. Colostrum is the first secretion of the mammary gland parturition and differs markedly from milk in its protein composition. I

^{*}From McMeekin, T. L. and Polis, B. D., "Milk Protein," Advances in Protein Chem., 5, 201-228 (

the first few days after the birth of the calf, the composition of the cal secretion gradually changes until that characteristic of milk appears. birth of the calf the colostrum contains approximately 17.5 per cent tein with about 5 per cent of this casein. The most striking difference the globulin fraction that carries the antibodies which give protection mst certain diseases. Since the health of the young animal often depends protection against some of these diseases, the importance of colostrum ling in both the calf and human infant has been studied. Euglobulin and udoglobulin are absent in the blood of the new born calf but when it is on colostrum they rise rapidly. Colostrum does not appear to be so intial in the feeding of human infants, since antibodies are more readily insferred by way of the placenta.

g Proteins⁸

The eggs of many species of birds have been eagerly sought and eaten by n. probably since the dim ages when he rummaged for food. The time of domestication of the chicken is unknown, but it was in the prehistoric rod. Whole egg is an excellent food because it is a very rich source not y of protein and lipid but also of most of the vitamins, except ascorbic J, and many of the required minerals except calcium. In some countries shells are used for food and are one of the major sources of calcium he diet. The proteins of egg are rather numerous and together yield an ellent assortment of amino acids for animals. They are considered to be he highest biological value.

he egg, of course, is produced for the nourishment of the embryo, ich develops in the fertilized egg. Some of the proteins of egg white e unusual properties which give them the ability to protect the embryo m bacterial invasion. Others possess properties with perhaps a regularly effect on the nutrition of the embryo. Lysozyme is antibiotic, ovocoid is a trypsin inhibitor, while ovomucin is an inhibitor of hemagtination. Avidin binds biotin, while conalbumin binds iron. The direct croons in the developing embryo have not as yet been demonstrated, but properties of these proteins are surely suggestive of how they may function.

he data considered here are those accumulated for chicken eggs. The set a few other species have had slight study; but when the term "egg" ed, it is meant to imply "chicken egg."

gg White Proteins. These have been studied for many years, and idreds of papers have been written about one or all of the proteins pressor far there is evidence for the occurrence of 8 different proteins in white. All have been isolated with a considerable degree of purity ex-

cept globulin G_2 and G_3 . Fevold⁸ prepared the data shown in Tab from existing data of a number of investigators.

Ovalbumin is the most abundant of the egg white proteins and is als protein studied most extensively. It is a phosphoprotein with a mole weight of approximately 45,000 and a small amount of carbohydrate. ious workers have reported that from 1.8 to 2.8 per cent of the mol consists of a polysaccharide composed of 2 glucosamines, 4 mannoses a nitrogen group. When the electrophoresis of crystalline ovalbum measured at various pH's, it is found that two peaks indicating two of molecules occur. On storage of the ovalbumin the relative proportion the two components changes. This has been determined on the basis of phosphorus content of the two albumins. One type of molecule is belift to contain two phosphoric acid residues, while the other, only one standing, some of the diphosphate probably changes to monophosph. Ovalbumin is readily denatured and precipitated by heat and a number denaturation reagents.

Conalbumin was recognized as early as 1900 by Osborne and Camp when it did not crystallize with ovalbumin. This is a protein wire molecular weight of approximately 70,000. It is able to bind iron in This property was discovered when egg white was found capable of rending iron unavailable to a microorganism, Shigelle dysenteriae, which quires iron in the nutrient medium. It was then discovered that althous the property was not lost on dialysis, it was lost on heating and eventual it was shown that conalbumin is the protein of egg white with this funct Ferric iron is bound to the protein molecule by coordination. Am groups, carboxyl groups, guanido groups, and amides are essential for activity of the protein. If these groups are blocked by reaction with sable reagents, conalbumin loses its power to bind ferric iron. The signals are selected as a selected as a

TABLE 4.4. PROTEIN COMPOSITION OF EGG WHITE*

Total protein	10-11 per cent wet basis
	82.8 per cent dry basis
Ovalbumin	70 per cent of total protein
Conalbumin	9
Ovomucoid	13
Lysozyme (G ₁)	2.6
(G_2))	
1	7
(G_3)	
Mucin	2
Avidin	0.06

^{*}From Fevold, H. L., "Egg Proteins," Advances in Protein Chem., 6, 90 (1951).

ce of the function of conalbumin for the development of the chick emois as yet unknown.

that it is not coagulated by heat. It has a molecular weight in the on of 27,000 to 29,000 and is a glycoprotein containing mannose and cosamine. It acts as an antienzyme for trypsin, diminishing the protease vity of the enzyme. The manner in which it is able to exert its innee has been studied. It has been shown that the groups on the ovocoid essential for antitrypsin activity are the carboxyl, guanidyl, and nolic groups and that they react with the amino groups on the trypsin, sino groups are not essential for the enzymatic activity of trypsin. Inition is consequently not the type where the antienzyme and substrate upete for a place on the enzyme, but instead inhibition represents some er change in the enzyme molecule.

he *globulins* of egg white consist of three different types of molecules ording to electrophoretic data. *Lysozyme* has received the greatest ount of attention because of its antibiotic activity. In 1922 Fleming blashed a paper on an agent capable of lysing or dissolving bacteria, ely distributed, and which he called "lysozyme." He found activity in saliva, plasma, numerous tissues, and a particularily active form in white. The activity of egg white has now been shown to reside in the fraction of the globulin. Since the antibiotic activity in this fraction is interesting, the name "lysozyme" rather than "globulin G₁" is widely 1. Lysozyme is a protein with a molecular weight of approximately 100 to 17,000. It is a basic protein with an unusually high percentage istidine, lysine, and arginine. Lysozyme is quite stable to heat, cold, many denaturation reagents. It is not stable in alkali. Some proteases has trypsin and papain do not attack it.

ilohulins G_2 and G_3 have received relatively little attention. Although G_2 been separated and shown to be a euglobulin, G_3 is still almost un-

wn as a separate protein. Neither have been crystallized.

Promucin is likewise relatively unknown because it is the most difficult the egg white proteins to handle, very insoluble, and consequently diffito purify. It precipitates from egg white on dilution with water at pH. The molecular weight is still in some doubt, but sedimentation data a value of 7,600,000. This interesting protein appears to be the one in this e capable of inhibiting hemagglutination. The ability of influenza to agglutinate red blood cells, called "hemagglutination," is inhibited egg white. It seems likely from studies of the activity of ovomucin tions that the hemagglutination inhibitor is ovomucin.

wilin is a protein present in low concentration in egg white, but it

nevertheless has been crystallized because of the interest in its ab bind biotin and render it unavailable to animals. If raw egg white in the ration to rats, they develop a biotin deficiency. Avidin is a pwith a molecular weight of not more than 70,000 as determined be mentation data, although other methods indicate a slightly lower values was the first naturally occurring substance shown to possess antivitantivity. The manner in which it renders biotin unavailable to an anima raw egg white is fed in the ration has not been discovered. It has shown that blocking most of the reactive groups on the protein mo amino, carboxyl, guanidyl and amide, and imidazole and phenolic not interfere with the activity. Some combination of protein and must occur, of course, but the nature of the bonds which hold the to the avidin are still obscure.

Egg Yolk. The proteins of egg yolk were first studied over 100 ago. Then followed a long period in which little attention was devo them. It is only in recent years that they have been reinvestigated. egg yolk is diluted with water, protein precipitates. When the yoheated, the proteins undergo heat denaturation and precipitate. Egg appears to contain at least two lipoproteins, lipovitellin and lipovite. Lipovitellin contains between 17 and 18 per cent lipid, while lipovite contains 36 to 41 per cent lipid, mainly lecithins. The protein portion these conjugated proteins are called vitellin and vitellenin, respect They can be prepared from the lipoproteins by exhaustive extraction 80 per cent alcohol. They are phosphoproteins and they appear to be similar except for the amount of phosphorus present. Vitellin con approximately 1 per cent phosphorus, while vitellenin contains only per cent.

Egg yolk also contains water soluble protein that does not precipital dilution of the yolk. This fraction is called *livetin*. It can be prepar precipitation on half saturation with ammonium sulfate, followe thorough extraction with alcohol-ether at -51°C to remove lipids. trophoretically there appear to be three components present in the frather enzymatic activity of the yolk is associated with this fraction.

The membranes within the shell and around the yolk are composed proteins which belong to the class of either keratins or mucins. Feedescribes how by staining techniques Moran and Hale indicate that the membrane is composed of an outer keratin layer, then two mucin followed by another membrane of a keratin and a mucin layer. The membrane appears by their method to consist of a mucin, keratin mucin membrane.

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The Flavor and Aroma of Food

FLAVOR IS THE SUBTLE and complex sensation that is the source of much the delight man finds in food. To both connoisseur and ordinary man, fla is of utmost importance in regulating preferences. It is the difference tween a cheap wine and the most expensive, the highest grade butter at the lowest, rancid cookies and fresh.

THE SENSATION OF FLAVOR

Flavor is a combination of taste, smell, and feel. In the mouth a pharynx are many taste buds capable of detecting sweet, sour, salty, a bitter. In the nose are olfactory endings that can detect a huge number different odors. The "mouth feel" of a food is likewise part of its flavor whether it is smooth or rough, tender or tough, unctuous so that it clings the tongue and roof of the mouth or watery so that it slides down read "Aftertaste" is a quality for the most part that is the combination of of these sensations after the particle of food has been swallowed. However, in aftertaste the sensation of feel has specific importance since stickiness or greasiness of the small amount remaining in the mouth and the teeth contributes to the general aftertaste.

Taste

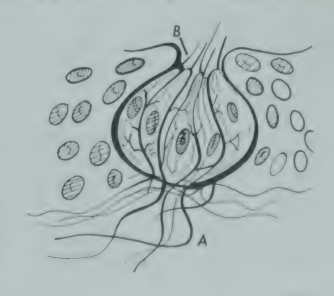
This sense is detected through the solution of soluble compounds in saliva or in the food juices and the contact of those dissolved compour with the taste buds. The commonly accepted theory (although not with challengers) of taste is that there are four primary tastes that can be tected: sour, sweet, salt, and bitter. The areas in which these tastes are tected overlap, but the sensation of sour is most readily detected on



FICURE 5.1. THE FLAVOR OF FOODS IS ONE OF THE DELIGHTS OF EATING. Orange, 10th, and vanilla are natural materials often added in small amounts to food for their or.

Courtesy of Food Materials Corp.

es of the tongue, salt on the sides and tip, sweet on the tip, bitter at the ck of the tongue and on the pharynx. The taste buds are present in great-tumber in the vallate papillae, the tiny nipple-shaped elevations disbuted in a V on the tongue. They are also present in papillae on the rest



TURE 5.2. A TASTE BUD. A. taste rs; B, taste pore. Reproduced from t, C. H. and Taylor, N. B., "The mg Body," Henry Holt and Co., New tk, N. Y., 1938.

of the tongue, the soft palate, pharynx, and epiglottis. The taste bud composed of a number of cells arranged in a tiny well around a nerve ing.

A compound to be tasted must occur in solution or dissolve in the same The solution seeps into the taste bud and the compound stimulate nerve ending. An impulse is transmitted along a nerve to the brain an recognize a taste. Then more saliva washes the solution out of the tiny

Taste thresholds are measures of the sensitivity to a given taste in to the detectable concentration. A common method for measuring threshold of, say, salt, is to give a number of samples with varying centrations of sodium chloride, a number of samples of pure water, sometimes of other compounds with other tastes. The subject attemp detect the taste present, rinsing the mouth after each attempt. The loconcentration consistent for an individual is his taste threshold for a ticular substance. The low levels at which tastes can be detected is a mazing. In a group of students almost everyone could detect 0.087 cent sodium chloride and 0.4 per cent sucrose. Considerable variation tween individuals occurs—a low threshold for one taste not always companied by a low threshold for the others.

Influence of Chemical Constitution. Taste depends on a number of tors, the most important of which is chemical constitution. Saltiness

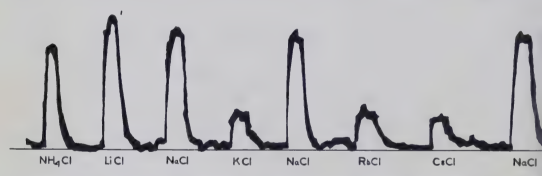


FIGURE 5.3. INTEGRATED ELECTRICAL RESPONSE OF CHORDA TYMPANI TO VARIOUS (CHLORIDE SOLUTIONS FLOWED OVER TONGUE OF RAT. Reproduced from Beidler, L. "Techniques and Methods for Research in Flavors, Chemistry of Natural Food Flavor A Symposium," Dept. of Army, Research and Development Command, Natl. Acad. Sci. Natl. Research Council, Washington, D. C., 1957.

property of electrolytes, and the halides particularily. In order of their s ness, the following ions are detectable: Cl, Br, I, SO_4 , NO_3 . cations also influence the taste. Na+ and Li+ salts tend to have salty to while K+, salts have considerable bitterness.

Sweetness is found in a number of organic compounds. Alcoholic droxyls tend to endow a compound with sweetness, but other groups sometimes effective. Saccharin, for example is a totally different c

and from sucrose, possessing no hydroxyl groups, but rather a sulfonnide. Sucaryl with 30 times the sweetness of sucrose is also a sulfoname.

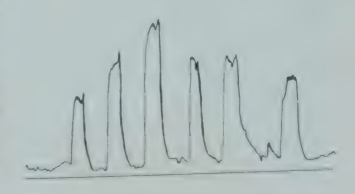
eneralizations concerning groups that make for sweetness in organic empounds are difficult.

Bitterness is a property of some organic and inorganic compounds. ome of the alkaloids such as quinine and brucine are exceedingly bitter, and NH₄, Mg⁻¹, and Ca⁻¹ are also bitter.

Sourness is a property of the hydrogen ion; its concentration is of priary importance in determining whether or not the sensation of sour is dected. The acids that occur in foods are organic with relatively low ionizators and consequently relatively low hydrogen ion concentrations. Added this is the common occurrence of a salt of the organic acid in the food, hich further lowers the ionization by common ion effect. Some workers lieve that aside from the effect on ionization, the anion likewise has an fluence on the perception of sourness. Research upon rats concerning eir nerve response to acids shows variation in response when hydrogen n concentration is constant. (See Figure 5.4.)

Influence of Other Factors. (Taste is also influenced by temperature, texre, and the presence of other compounds. Mackey and Valassi¹⁴ meased the taste thresholds for the four primary tastes in water as well as to mato juice and in egg-milk custard prepared as liquid, gel, and foam, hey found that the primary tastes were harder to detect in gels than in quids and that the foams were intermediate.

FIGURE 5.4. RESPONSE OF RAT TO RIOUS ACIDS. The large peaks m left to right show response to drochloric, citric, formic, oxalic, condend from Beidler, L. M., "Techtees and Methods for Research Mayors. Chemistry of Natural cod Flavors – a Symposium," Adv. ard on Quartermaster Research Development, Dept. of the Army. 15hington, D. C., 1957.



The effect of one primary taste on another is well known to ever who cooks. Salt tends to decrease the sweetness of sucrose, so that to which salt is added has a richer, less sweet taste than that without Sugar likewise can tone down and round out saltiness. Many a consalvaged the mashed potatoes that were too salty, by adding a amount of sugar. The effect of sweet on sour, such as sugar on lemo or vinegar, is also important. Any food quite sour and yet quite sweat rich, full flavor that is lacking when either sourness or sweet present alone. Salt also tones down sourness of food acids. This down of sensation when two tastes are presented simultaneously is compensation.

Successive contrast, on the other hand, tends to sharpen a sense Grapefruit seems unusually sour if eaten immediately after a sweet A plum tastes quite sour after candy but quite sweet after grapefruit. An interesting phenomenon has been discovered in relation to the of a few compounds. Sodium benzoate tastes quite different to differe dividuals and the threshold at which it can be detected is very differently thiocarbamide (PTC) has been extensively studied: "taste ness" to this compound is inherited according to Mendelian ratio. It parents are nontasters, the children do not taste PTC. Women tend to a slightly lower threshold than men do for this compound. Taste blin in women occurs in 22.2 per cent and in men 25.9 per cent.

Fatigue for taste does not occur rapidly and some physiologists delieve it ever occurs. However, it is a common experience to no diminution in the taste of a food as one continues to eat it. Some periments indicate that fatigue occurs for one taste but the others are

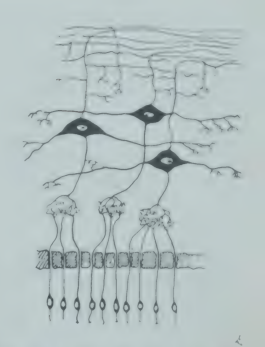
affected or are even enhanced.

Odor

We can detect differences in the odor of thousands of compound pinging on the olfactory nerve endings. The sensation is experience course, only when the nerve impulse is transmitted to the brain are corded there. Our remarkable ability to remember odors and the ability the brain to receive and detect differences in odors cannot be adequively explained at present. Many people can recognize odors that they have perienced only once, many years before. The author once heard a 70 old woman proclaim that a noxious odor must be a stinkhorn must since she remembered smelling one as a girl. The hoots of derision quieted when a stinkhorn was found to be the source of the odor. individuals have experienced this remarkable recall for odors.

The aroma of food is recognized as a tantalizing and delectable pro-

mom are cells of olfactory mucous membrane has nerve fibers pierce skull and reach procof olfactory bulb cells. Reproduced from C. H. and Taylor, H. B., "The Living N." Henry Holt and Co., New York, N. Y.



tating but not always as an invaluable part of flavor. We have only to affer loss of our ability to detect odor through congestion of the nose, to falize what a contribution it makes. Food tastes salty, sweet, sour, and after when we have a bad cold, but how flat and uninspired it is without dor. All the subtle nuances of the flavor complex are gone.

The olfactory mucous membrane is located in the upper part of the nasal with on portions of the turbinates and septum, and contains olfactory res imbedded in a special epithelium. Glands are present that secrete aid. In order for a chemical compound to possess an odor it must vapore and pass into the nasal cavity. It is not known whether the molecules ssolve in the fluid on the lining and the solution comes in contact with e olfactory cells or whether the hairs of the cells penetrate the mucous ic come in contact with gas. The nerve then transmits an impulse to the ain. Allison and Warwick have shown that a rabbit has about 10.000,000 receptors, each with 6 to 12 hairs. This is a tremendous area r receiving odor stimuli. The detection of an odor during breathing ocirs through the diffusion of the gas into the still air of the upper cavity. re olfactory endings are not located on the epithelium in the direct path the inspired air. When we detect a flavor in a mouthful of food, the formus compounds change to gases and diffuse through the pharynx into e. As we raise the food to our mouth, we inhale its odor.

The sense of smell varies considerably between individuals and is, of surse, much less keen in man than in many other animals. Nevertheless, is still remarkably keen for some compounds. Vanillin can be detected most individuals at a concentration of one part in ten million.

Fatigue for odors occurs very quickly. We have all observed how st an odor of a food such as coffee or meat will be when we first come in building or room where it is being prepared and how quickly we loose sensation. Students who do not work in chemistry, often complain o odors of the laboratory while those of us who work there constantly not detect them unless the concentration suddenly increases or a new g formed. People have been overcome by hydrogen sulfide, which has an usually strong odor, after fatigue of the olfactory sense was so depre that they forgot the concentration was high.

Although fatigue for odors occurs rapidly, it is selective. We may no able to smell the odor of a compound after a short period of time, but sense of smell for another odor is unimpaired.

Testing Sense of Smell or Aroma. Elsberg⁶ devised a method for test acuity of smell. A flask is filled with cotton and between the layers toothpick, cotton wrapped, that has been dipped in a solution of some sential oil or odorant, often a pure chemical. Air is admitted to the fin measured amounts and the vapors sniffed. The amount of air requires perceive the odor and also the concentration of the solution on the congive a measure of the acuity of the individual to that particular odor. So individuals have high acuity of smell but cannot readily recognize od Thus, another part of the test often introduced is proper description of odor perceived.

Some physiologists are now studying taste and odor responses in animal and attempting to obtain objective data that will make possible an unstanding of these phenomena. Since the impulse that travels over a nervelectrical, measurement of the potential of a nerve shows the effect of stimulus. It is possible in a living animal to make measurements on large nerve bundle that serves many taste bads, on an isolated fiber shing the response of several taste cells in one taste bud, or by insermicroelectrodes to make measurements on a single taste cell. Olfactory sponse is more difficult to measure although record of the activity of olfactory nerve can be made. It is also possible to measure the change potential in the presence of an odorant across olfactory tissue after excised from the animal. None of these studies is easy, but it is here we can hope to find better understanding of the physiology of taste odor. 15

Feeling

A Some compounds not only stimulate the olfactory ending but also that the trigeminal nerve—a nerve with endings in the skin of the face, ton and teeth, giving a general sensation. Ammonia, a common substance is

mulates this nerve, has an odor for us but also another sensation deibed as "tingling" or "sharp."

Other "feel" factors contributing to the over-all sensation of flavor are hot, burning effect of some peppers and spices as well as the coolness peppermint. The texture of the food is also important, grainy fudge tes sweeter but not as rich as creamy fudge. Crispness and stickiness other sensations of mouth feel. Some wines contribute to flavor by ming to fill the mouth and cover the pharynx immediately after a small

ends

A blend of sensations occurs with most foods so that sorting out all of em is difficult. This is often highly important to the satisfaction that a perbly delectable dish affords. Many foods have a slightly sour note that est people do not separate as "sourness" but simply as part of the dehtful flavor. Broccoli, for example, has a volatile acid which is part of fine flavor and most honey contains appreciable amounts of acid. Ocsonally elements of flavor appropriate and desirable in some foods are thly repugnant when present in others. For example beer has a sour. rsev aroma and some ripened cheeses have an element of rancidity in the or but, because it is expected and blended, it is pleasing. In the Arthur the Flavor Laboratory scientists have tried blends of many ingredients, re compounds and extracts, and conclude that in blending various reacns may result. With two odorants: (1) some of the major notes of each trant may be suppressed; (2) all of the major notes of each odorant may suppressed: (3) some of the major notes of one odorant are suppressed trone of those of the other odorant: (4) a complete blend takes place th the formation of a new odor; or (5) a partial blend occurs with a new or formed but some characteristics of each odorant retained. There is Il much to learn about blending, masking, and enhancement when two or re odorous compounds are mixed.

Appropriateness is of great importance in our enjoyment of flavors in ods. An onion flavor may be delectable in a stew or soup but objection-le in a custard. We become conditioned to expect certain sensations from tain foods and while a slight variation is titillating, a completely unsected taste is unacceptable. Most of us have a "sweet tooth" and relish ing sweet. But while we may find that chicken prepared with the adon of a small amount of thyme is exceedingly interesting because the cor departs a little from that which we usually experience, chicken presed with a strongly sweet sauce may be rejected as improperly prepared

d unappetizing.

Aftertaste is part of the sensation of flavor of some foods. It is ticularily common with foods that leave a residue in the mouth swallowing. Sticky foods such as syrup or greasy foods such as meatend to leave a residue coating the mouth for a short time after swalling and all or part of the sensations which are part of flavor, controbe experienced. The persistence of the flavor is highly desirable in sfoods, although usually we do not enjoy the sensation if it persists long. Many people do not eat raw onions because of the persistent a taste. The volatile disulfides present are absorbed by the blood and then secreted in the saliva and exhaled in the breath, so that the odor persist for many hours.

CONTROL OF FLAVOR AND AROMA IN PROCESSED FOOD

Control of flavor and aroma in processed food is of utmost importa in determining the quality and selling price of the finished product; is likewise true of simple cookery operations. Many factors, all of w must be considered, influence the flavor. The quality of the ingredients of course, a considerable effect on the product. Off-flavor ingredients not produce an item with excellent flavor. Conditions of processing mus carefully controlled. Thus in roasting coffee or cocoa beans the temp ture and length of time roasted must be controlled in order to insure fla of the right quality. Avoidance of contamination by flavorful compou during processing or storage must be watched. Thus the lacquer lining cans cannot contain soluble flavored substances. Many foods such as f and baked goods, pick up odors of other foods readily and must be a quately protected from contact with these odors. Foods must also be p tected from contamination by bacteria and molds and stored under of ditions where these microorganisms cannot grow rapidly. Molds prod compounds that impart a musty or sour flavor which is completely un

In many food industries, tasters are employed to grade the product often the raw materials. There are usually no chemical tests that can ferentiate the subtle differences between a quality product, an ordinary and one that is off-flavored. With a little experience, some tasters deveremarkable ability to differentiate products. Some can readily detect flin the processing operation and spot the location of the trouble by taking many industries these tasters are of great value to the company states one of the bases on which the consumer buys the product.

Good flavor in a food product is desired by everyone who works in figroduction whether a housewife or the director of a nation-wide indus

ul³ has analyzed the pattern of good flavor as the following sensations: ran early impact of appropriate flavor; (2) rapid development of an imston of highly blended and usually full-bodied flavor; (3) pleasant the sensations; (4) absence of isolated unpleasant notes; and (5) anticiton of the next mouthful."

asurement of Flavor

ine sensations of flavor are such complex reactions that it is impossible my simple chemical or physical test to measure them. An appreciation avor depends on a human being and his reactions to the actual act of ing. Since the flavor of food products is so important in determining a commercial value, many methods for measuring flavor have depend. These have been reviewed at various times. A recent review of methods and their implications is in a symposium in Food Technology 1957.

xpert tasters are employed in some industries, particularly for wine, iskey, tea and coffee, and spices. The expert taster has usually grown up are industry and is of value only to this industry. Through interest and cortunity he has developed a "discriminating palate" for the food of company. There is no indication that an expert taster has a keener se of taste than the normal individual, but rather that he has years of ning.

Other methods use a panel of tasters; sometimes these are individuals in some training, sometimes they are workers in the industry with sufficient interest to be willing to serve on a panel, and occasionally where terence is desired, they are large numbers of consumers. The panel may icate its preferences or judgment of quality by scoring a food on some laefined qualities. Often a numerical scale is used so that the scoring he individuals can be added readily to give a composite score. Scoring been widely used in meat, bakery, and milk products and in recent rs with dehydrated foods.

Piperence tests are sometimes used in an attempt to get a more precise reproducible test of flavor in foods. A trained panel is required for se tests. Boggs and Hansen have summarized the various methods of erence testing. Either a pair of samples or a triangle in which two of samples are identical and the third is to be separated are presented be judges. Sometimes in a triangle a standard is submitted and one of a is matched to it. In other tests the judges only know that two samples identical.

he dilution test is a type of difference testing. A sample is presented to judges and then other samples that may or may not contain the unown at a definite level of dilution are offered. The method was first applied to the detection of dried egg in fresh egg. The higher the of the dried egg the greater the difficulty of detecting it. In order to high degree of agreement, it would be necessary to have a high pedried egg present. A poor quality product would be detectable at a centration or a high dilution.

Ranking is used with a flavor panel and with either a numerical betical evaluation of one property. A series of samples are supplie member of the panel and he arranges them in the order of increasi creasing quality of the characteristic. A high degree of agreement members of a panel occurs when the samples differ to a marked dewhen differences are subtle, agreement is not as common.

The flavor profile is a method for evaluating a flavor by desc either as a whole or by characteristics. The vocabulary by wh characteristics are described are comparative terms such as "rubb "eggy;" or, when possible, they are referred to a compound—pher acid for "horsey." Feeling effects are described on the basis of th throat or nose reaction, such as "throat burn," "puckering," an ing." The reactions are broken down into (1) character notes, (2) appearance, (3) aftertaste, and (4) amplitude. The character notes protruding sensations, often exceedingly difficult to distinguish in The order of appearance may appear at first sight to be of little im or even absent. But close attention to flavor during tasting will re many flavors are not perceived as a whole but rather as a series tions. These come very rapidly but can be separated. Coffee bitt not perceived until after the other flavors have been experienced. an unpleasant note should not be either the first or last sensation flavor. Aftertaste has been mentioned before, and its importance taste complex is readily recognized. "Amplitude" is the term use press the total effect of flavor and aroma. The fitness of the fla broad aspects of all characteristics is included here. It requires s derstanding and experience to judge amplitude. It is rated as very 1 medium or high. Examples of amplitude can show the meaning term. Tomatoes picked green have low amplitude, those grown house are medium, and the vine-ripened are high. Potatoes, boiled seasoned, have low amplitude; salt added at the table improves the tude by blending the flavor but salt added in preparation gives hig tude.

Any evaluation by a flavor panel is subject to error. Many of cautions necessary to the satisfactory use of a panel have been and described. It is essential that the members work alone in quie surroundings. Usually air-conditioned booths are used so that



© RI 56. OFFACTORIUM FOR TESTING ODORS, Courtesy of Food Acceptance Branch, attermaster Food and Container Institute for Armed Forces, Chicago 9, III

perature and the humidity can be controlled. To avoid influencing one ther, the panel members do not discuss their reactions with each other, in the day of the week and the hour of the day has an effect. Mitchell 13 that the sensitivity of a duo-trio test where two samples out of three packed, is greatest on Tuesday with Friday also better than Monday. The day, or Thursday. In an eight hour day accuracy of judgement was need during the fourth, fifth, and sixth hours.

p schological factors, the environment must be carefully controlled are ser, some of the psychological factors that cannot be controlled are

the experience background of each tester and his personal problem reactions. Food experience affects everyone. If you have frequently food highly seasoned with garlic, you will usually like that flavor but have nothing of newness or surprise for you. A person adventurous in life experiences will find pleasure in a new eating experience, while the conservative will dislike anything unfamiliar.

The consumer acceptance or preference has been used in recent year some products. If a new food is introduced, only one sample is offere large consumer panel, but if a food is modified, then two samples are mitted and a preference requested. Occasionally more than two samples submitted but usually this leads to confusion in the subjects and the rare doubtful.

Flavor Intensifier: Monosodium Glutamate

Large quantities of monosodium glutamate have been used in the of for years and in the United States for approximately 2 decades as a intensifier. In China and Japan fermented soybean curd is added to foods and soy sauce is a common condiment. Soy sauce is rich in a sodium glutamate and interest led eventually to the production of atively pure monosodium glutamate. It is claimed by some authors that compound has little or no flavor itself but intensifies the flavor of and vegetables through a rounding or blending effect. It is widely us stews, canned meats, soups, chowders, etc. It does not have an effect fruits or fruit juices or sweet spicy foods. Cairncross and Sjöström ported that it suppresses the sharpness of onion, the rawness of many tables, and the earthiness of potatoes. They also reported that the effinoticeable in the pH range 3.5 to 7.2, the range in which most food eaten. They found that fats, oils, and high viscosity in foods modifinfluence of sodium glutamate considerably.

The mode of action of sodium glutamate and its effect on flavor are in dispute. Much of the work carried out has been an investigation of preference of tasters for a food with and without added monosof glutamate, and the underlying physiology and biochemistry of the effect of the effect of the control of

Glutamic acid occurs in high concentration in numerous proteins. We gluten for example contains approximately 36 per cent. The free glut acid hydrochloride is prepared commercially from wheat, corn or separet wastes by acid hydrolysis of the protein. Glutamic acid is formed dissolving the crude hydrochloride, adjusting the pH to 3.2, and allow to crystallize slowly. The product is neutralized with either sodium droxide or sodium carbonate, decolorized and crystallized. The product relatively pure monosodium glutamate

Very large amounts of this material are now used commercially in the oduction of many meat and vegetable dishes. Although it has been smed that it is not detectable and although it is true that pure monodium glutamate is only slightly sweet and salty, the author has noticed at the presence of this "intensifier" at the level usually used in foods 1 be recognized. It gives a sameness of flavor to foods as dissimilar as eken or tuna pies and makes the product markedly different from the shly prepared dish. On this basis, it is highly objectionable to her.

avoring Extracts

These are widely used in the food industry, and in the United States quite well standardized. The Food and Drug Administration defines an ract as "a solution in ethyl alcohol of proper strength of the sapid 1 odorous principles derived from an aromatic plant, or parts of the nt, with or without its coloring matter, conforming in name to the plant 2d in its preparation." If artificial coloring or synthetic flavoring computed are added, it is required that the label state this. The flavoring exects are made either by adding the essential oil to alcohol or to water dalcohol, or by percolating the chopped plant or plant part with a mixtee of water and alcohol. Thus lemon extract is prepared by adding the non oil expressed mechanically from lemons, to alcohol or diluted alcohon oil expressed mechanically from lemons, to alcohol or diluted alcohon oil expressed mechanically from lemons, to alcohol or diluted alcohon oil expressed mechanically from lemons, to alcohol or diluted alcohon oil expressed mechanically from lemons, to alcohol or diluted alcohon oil expressed mechanically from lemons, to alcohol or diluted alcohon oil expressed mechanically from lemons, to alcohol or diluted alcohon oil expressed mechanically from lemons, to alcohol or diluted alcohon oil expressed mechanically from lemons, to alcohol or diluted alcohon oil expressed mechanically from lemons, to alcohol or diluted alcohon oil expressed mechanically from lemons, to alcohol or diluted alcohon oil expressed mechanically from lemons, to alcohol or diluted alcohon oil expressed mechanically from lemons, to alcohol or diluted alcohon oil expressed mechanically from lemons, to alcohol or diluted alcohon oil expressed mechanically from lemons, to alcohol or diluted alcohon oil expressed mechanically from lemons are expre

Yanilla extract, the most widely used extract in the United States, is pred by percolating macerated vanilla beans with alcohol, frequently in presence of glycerol or sucrose. The flavorful compounds extracted are crous but the most abundant is vanillin. Since vanillin can be preced synthetically very cheaply, synthetic vanilla is sold in large quantities. If ever is not quite as fully rounded as real vanilla extract but it is used tely in many foods selling at relatively low cost. Usually coumarin is

Vanillin

beans. Discussion of the methods of preparing flavoring extracts c

added to the vanillin in the synthetic extract to improve the flavor, an amel or a food dye to simulate the brown color extracted from the v

found in a textbook of food technology.

Synthetic Flavoring Substances

These have been produced for many years. Originally esters were usuall amounts for imitation flavors, particularly in cheap confection though an ester such as amyl acetate is reminiscent of banana flavor, sesters are crude, rough substitutes for natural flavor. Analysis of national flavors has developed slowly because of the difficulty of separating stances that occur in a concentration as low as that in most foods, methods of study (see below) are now opening the way for careful an tailed analysis of the flavoring compounds in many foods. During the 50 years numerous compounds have been synthesized which are use synthetic or imitation flavors. The common so-called *flavormatics* are in a table prepared by the Food Additive Committee of the Flavorin tract Manufacturers Association of the United States. (See Table These are arranged in order of their importance. The type of flav which they are usually used and the parts per million concentration if finished food are given.

Table 5.2 shows the composition of a few imitation flavors. It is inting because it shows the complex nature of some of these imitation flat The list of compounds isolated from natural flavors is far from com Many more compounds have now been identified in strawberry, apple coffee flavors.

RECENT DEVELOPMENTS IN FLAVORING RESEARCH

Recent developments in flavoring research have followed three prir paths: (1) careful identification of the composition of some flavors, ticularly through the use of gas chromatography as a means of separ

(text continued o

TABLE 5.1. THE PRIMARY FLAVORMATICS

THE PROPERTY OF THE PROPERTY O			
Aromatic Chemical	Flavor	Avg. Use (P.P.M.)	B.P.°C
oup I			
nillin	vanilla	31.5	81.5° MP
hyl Vanillin	vanilla	16.6	76.5° MP
ral	lemon	17.6	229°
nzaldehyde	cherry, almond	84.8	180°
namie Aldehyde	cinnamon, cola	110.7	252°
ethyl Salicylate	root beer wintergreen	129.7	221°
nyl Acetate	banana, fruit flavors	78.4	142°
nyl Butyrate	banana, fruit flavors	23.7	185°
hyl Acetate	fruit, rum	92.7	76°
hyl Butyrate	strawberry, fruit	31.1	120°
ethyl Anthranilate	grape	97.1	255°
hyl Oxy Hydrate	rum	472.7	
ethole	root beer, anise	133.5	235.3°
frol	root beer, anise	16.9	236°
enthol	mint	111.2	215°
oup II			
actropine	vanilla, cherry	3.4	263°
denvde C-18	coconut	17.6	263°
scetvi	butter	17.3	88°
denvde C-16	strawberry	10.0	271°
n Oenanthate	grape	8.1	187°
VI Caproate	pineapple	11.7	187°
Jehyde C-14	peach	8.0	297°
ger.ol	spice	48.8	253°
nvi Valerinate	peach, fruit	9.6	200°
w Valerinate	fruit	14.4	145°
rvene	spice, mint	190.3	230°
is Aldehyde	cherry, vanilla	3.6	247°
oup III guo			
ral Acetate	strawberry, fruit	8.8	215°
wl Lormate	rum, strawberry	77.0	54°
)! 3	raspberry	0.9	147.5°
vilactate	grape	42.1	154°
Ivl Videhyde	cherry	9.3	204°
n caproate	pineapple, apple	4.4	222.2°
lous	spice	30.2	199°
menal	rose (general)	14.20	206°
** Acetate	fruit	26.4	126.5°
Al Petargonate	grape, fruit	2.3	- 228°
W Barvrate	fruit	17.9	166.4° 208°
(e) (de (-10)	citrus	1.2	208

TABLE 5.1 (Continued)

Aromatic Chemical	Flavor	Avg. Use (P.P.M.)	B.P.°C
Ethyl Heptoate	grape, pineapple	4.6	172.1°
Geranyl Acetate	fruit	8.4	245°
Methyl Phenyl Acetate	honey	2.4	222°
Group IV			
Aldehyde C-8	citrus	0.71	165°
Aldehyde C-9	citrus	2.81	191°
Cyclohexyl Butyrate	pineapple, fruit	8.3	212°
Ethyl Laurate	spice, fruit	6.9	269°
Linalyl Acetate	citrus	5.4	220°

^{*}From Janovsky, H. L., Food Technol., 9, 500–502 (1955).

TABLE 5.2. SYNTHETICS*

	Aromatic Isolated	Sample Formula	
Raspberry	beta ionone anisic aldehyde benzaldehyde phenyl ethyl alcohol hexene n-hexanol iso-amyl alcohol iso-butyl alcohol	beta ionone iso-butyl acetate anisic aldehyde phenyl ethyl alcohol phenyl ethyl iso-butyrate ethyl methyl p-tolyl glycidate phenyl ethyl anthranilate vanillin hexyl butyrate iso-amyl acetate ethyl n-butyrate rose oil diallyl sulfide	3
Strawberry	borneol methyl caproate amyl aldehyde ethyl caproate	ethyl methyl phenyl glycidate amyl aldehyde bornyl acetate ethyl caproate vanillin beta ionone ethyl methyl p-tolyl glycidate iso-butyl acetate ethyl butyrate	1 3
Cherry	geranyl butyrate ethyl caprate terpenyl butyrate ethyl caproate benzaldehyde geraniol	cinnamyl anthranilate iso-amyl acetate iso-butyl acetate cinnamic aldehyde dimethyl acetal benzyl alcohol geranyl butyrate ethyl caprate	1

B.E 5.2 (Continued)

	Aromatic Isolated	Sample Formula	
-			7-
-17)	63	terpenyl butyrate	0.5
ntinued)		ethyl caproate	2
		benzaldehyde	8
		p-tolyl aldehyde	3
		p-methyl benzyl acetate	11
		ethyl methyl p-tolyl glycidate	16
		vanillin	7
		heliotropin	5
ich	linalyl esters (as	linalyl formate	10
	valerate, acetate,	gamma undeca lactone	13
	formate)	linalyl butyrate	4
	furfural	heliotropin	14.5
		geranyl valerate	15
		furfural	1.5
		alpha-methyl furyl acrolein	6
		methyl cyclo pentenolone valerate	10
		benzaldehyde	5
		iso-amyl formate	15
		iso-butyl butyrate	6
pie	geraniol	geranyl valerate	10
	acetaldehyde	geranyl n-butyrate	8
		geranyl propionate	8
		linalyl formate	10
		iso-amyl valerate	15
		vanillin	8
		allyl caprylate	6
		geranyl aldehyde (citral)	5
		acetaldehyde	6.5
		methyl cyclo pentenolone valerate	8
		alpha methyl furyl acrolein	2
		iso-amyl butyrate	13.5
	α -furfuryl mercaptan	α -furfuryl mercaptan	1()
.10.	(2-turiary) mercaptan	ethyl vanillin	3
		solvent	87

From Katz, A., Food Technol.. 9, 636-638 (1955).

targe number of compounds that occur in low concentration in many trail foods, (2) production of natural essences by condensation of the pors formed in vacuum or low temperature concentration of foods, and the search for flavor precursors and their enzymes. As the food instry moves farther along the path of prepared or partially prepared of the problem of flavor in the final product becomes more trouble-

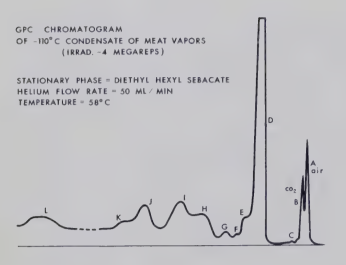


FIGURE 5.7. GAS CH GRAM OF MEAT VAPORS. pip for air, B for CO₂, subsequent rises are vario pounds present in the var produced from Pilgrii Schutz, "The Chemistry of ral Food Flavor—a Symp Dept. of Army, Research velopment Command, Nat Sci. and Natl. Research Washington, D. C., 1957.

some. With the concentration of populations in cities and the rise in ard of living, a growing demand for quality food occurs at the same as a need for food products stable enough to allow time for distrib Maintenance of flavor is one of the chief problems in producing q food.

The gas chromatograph is a fine instrument for the separation of many compounds that make up the flavor complex in a natural food. The packed with finely divided particles of a solid, and wet with a of low volatility. The nature of the liquid affects the separation ach and as in all types of chromatography, the discovery of the suitable is often one of the most time consuming parts of the research. A contrate of the essence is made by one of a number of methods—for exact extraction or distillation—and this concentrate is placed on the contrate column in fractions or as single compounds. The fractions are reported to the columns and then separated into single compounds. Separation very small quantities of material can be made very rapidly in a material minutes and the recoveries from the columns are almost quantitative. Large samples the gas chromatograph can be used as a preparative device.

After the components of a mixture are separated, the classical me of identifying compounds can be used. In regular paper or column chr tography, the identity of a compound is often established by compa of its behavior with that of a known compound. This technique may be plied to gas chromatography. Relative amounts of substances are reco

The flavors of many foods are now being studied by this method and fairly short time the chemistry of flavor will be on a firm foundation the first time. An understanding of changes in flavor on processing or of a food such as coffee is at last possible with this technique. Tables 5.3. and 5.4.)

TABLE 5.3. COMPOUNDS FOUND IN THE GASEOUS EMANATION OF ONIONS

Relative Amount	
very abundant	
abundant	
small	
trace	

During the processing of a number of foods, particularly fruit juices, loss of aroma during concentrating is appreciable. Many years ago ruum evaporation was introduced with low temperature condensors to as much of the volatile compounds as possible. This was a great important in many cases over evaporation at atmospheric pressure. However, since air is present in the original product, venting the system is cessary; consequently, enough volatile material is lost so that the fresh

TABLE 5.4. VOLATILE COMPONENTS IN COFFEE AROMA

Formic Acid	Phenol
Acetic Acid	Resorcinol
Methyl Ethyl Acetic Acid	Cresols
n-Valeric Acid	Ammonia
iso-Valeric Acids	Methyl Amine
Higher Fatty Acids	Trimethylamine
Ethyl Alcohol	Pyrrole
Acetyl Methyl Carbinol	n-Methyl Pyrrole
	Pyridine and Homologues
Furfuryl Alcohol	Pyrazine
Acetaldehyde	Guaiacol
Methyl Ethyl Acetaldehyde	p-Vinyl Guaiacol
Furfural	Eugenol
Acetone	Hydrogen Sulfide
Diethyl Ketone	Methyl Mercaptan
Diacetyl	Dimethyl Sulfide
Acetyl Propionyl	Furfuryl Mercaptan
Hydroquinone	
Esters	Furane
Furfuryl Formate	Sylvestrine
Furfuryl Acetate	Vanillone
Methyl Alcohol	n-Heptacosane
2,3-Dioxyacetophenone	

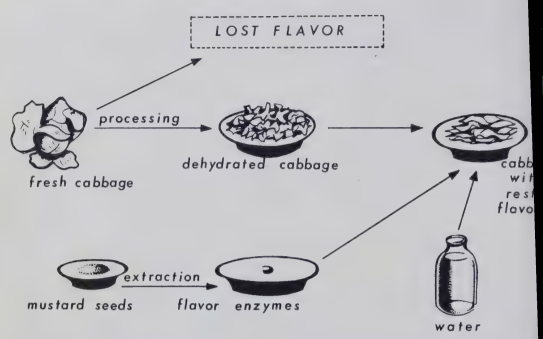


FIGURE 5.8. RESTORATION OF NATURAL FLAVOR TO DEHYDRATED CABBAGE. R duced from Hewitt, E. J., MacKay, D. A. M., Konigsbacher, K. S. and Hasselstrom "Flavor Propagation Through Enzymatic Action, Chemistry of Natural Food Flavor symposium," Dept. of Army, Research and Development Command, Natl. Acad. Natl. Research Council, Washington, D. C., 1957.

flavor disappears. Improvements in design now allow the remova most aroma in the first fraction by flash evaporation. The juice is he rapidly in a matter of a few seconds by passing it with turbulent through a small pipe and then it is rapidly cooled. Flash evaporatio carried out at atmospheric pressure if the volatiles will take the temperature, otherwise at low pressure. The condensate from the fevaporation passes to a fractionating column and is separated into fractions. The vent gases are scrubbed by a counter-current of cold column that trap the most volatile compounds by dissolving them. The sence is stripped from the column.

With the progress of biochemistry and the rapid rise in our understating of the changes that occur in plant and animal cells, the time is for some investigations of flavor precursors and attempts to utilize chemical reactions for the maintenance and enhancement of fresh flated Studies are underway to find methods for the isolation or at least the centration of flavor precursors and enzyme systems capable of developing fresh flavor. It is hoped that when these are obtained their additional dried or concentrated foods will improve the flavor and acceptability of in the presence of off-flavors.

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IAPTER SIX

leat and leat Products

r is defined by the Food and Drug Administration as follows: "Meat properly dressed flesh derived from cattle, from swine, from sheep or sufficiently mature and in good health at the time of slaughter, but is reted to that part of the striated muscle which is skeletal or that which and in the tongue, in the diaphragm, in the heart, or in the esophagus, does not include that found in the lips, in the snout, or in the ears, or without the accompanying and overlying fat, and the portions of sinew, nerve, and blood vessels which normally accompany the flesh which may not have been separated from it in the process of dressing r sale.

lesh is any edible part of the striated muscle of an animal. The term all as herein used, indicates a mammal, a fowl, a fish, a crustacean, a usk, or any other animal used as a source of food."

MAL STRUCTURE

scle

here are three types of muscles in animals: striated or voluntary muscle the constitutes the meat in the above definition, smooth or involuntary cle which for the most part is discarded when an animal is dressed, be intermediate between the other two muscles.

riated muscle is composed of long cylindrical cells, called the muscle s, which lie parallel to one another and lengthwise of the muscle. Each contains a number of nuclei lying close to the outer edge near the ibrane or sarcolemma. The cross striations that appear under the

microscope have been the object of numerous studies but as yet a co explanation of their appearance has not been established. The musc are bound together by connective tissue and groups of them are asso and covered with more connective tissue (the *perimysium*) to form but In some muscles the bundles are very noticeable when the muscle crosswise, giving the grain to the meat. The bundles of muscle fibe held together in a single muscle by a covering of connective tissue called the *epimysium*.

The cells of the muscle fibers contain a complex mixture of prolipids, carbohydrate, salts, and other compounds. The carbohydrate ing muscle is glycogen, but during slaughter and the ripening that between the killing of the animal and the cooking of the meat, the gly is degraded and disappears from the tissues.

Muscle also contains lipid—phospholipid and cholesterol. However termining how much lipid is in the muscle cell and how much in the nective tissue is difficult. Striated muscle is believed to contain ab per cent lipid. The connective tissue contains large amounts of neutre that marbles the meat and is distributed between the bundles of medical cells and is of great importance to meat quality at the table. It is distoremove lipids completely from the muscle cells.

The proteins of a muscle cell are numerous and comprise those impo in contraction, in the functions of the nucleus, and in the enzymatic tions of the cell. These latter may not be completely different from others—for example, the protein myosin is necessary for contraction also as an enzyme for ATP (adenosine triphosphate). The classical me of protein fractionation by extraction with salt solutions has been ap to muscle. This method often leads to fictitious separations and some of older work is consequently open to question. Two proteins present i cells have received much attention in recent years because they are sponsible for the contraction of the muscle fiber. The two protein actin and myosin and the compound formed between the two, actomy A protein fraction can be extracted from muscle with cold water. fraction is called myogen and comprises a number of proteins incli those enzymes essential for glycolysis. Many of the enzymes promotin long series of reactions that occur when glycogen undergoes fragment have been isolated, purified, and some have even been crystallized. pigment responsible for the pinkish red color of muscle, myoglobin also received some attention and is more completely discussed on p. Another group, the proteins called nucleoproteins, is now being st extensively.

he state of our knowledge of the proteins of a muscle cell is similar to to our knowledge of any cell. Biochemistry is at the state where technes for the separation of cellular compounds and an understanding of role at the level of the cell are still developing. In the past it has possible to fractionate tissues crudely, although separation of proteins pure compounds has been uncertain. Now not only are techniques and wledge growing to the point where criteria of purity for large molecules as proteins are known, but also methods are available for separating sof cells such as nuclei, mitochondria, and cell membranes.

ne salts in muscle cells are the same as those in most cells of the body are at the same concentration. Potassium is the most common cation, i magnesium and sodium following. The anions are acid phosphate, rbonate, and sulfate in the order of their concentration. The contrations of these ions cannot be determined directly because the cells in contact with extracellular (interstitial) fluid, and at present it is not sible to separate cells completely from extracellular fluid. In meat we k relatively large pieces of tissue and consequently have both extracellular dintracellular fluid. The salts in the extracellular fluid have a contration equal to that in the blood plasma. Sodium is the most important on with potassium, magnesium, and calcium in small amounts. The stabundant anion is chloride, with fair amounts of bicarbonate and ll amounts of acid phosphate and sulfate.

he organic compounds that can be separated from muscle by treatment water and that are not lipid or protein are called the extractives. These stances appear in the water when meat is stewed, fricasseed, or in any treated with moist heat and in the pan juice when meat is roasted or 1. Striated muscle has approximately 1 per cent organic extractives and r cent salts. Glycogen is the most abundant extractive in resting muscle, as pointed out above, the amount decreases after slaughter. None can etected in most cuts of meat on ripening. In the extract are products lycolysis or intermediates from the tri-carboxylic acid evele such as ic acid and pyruvic acid and small concentrations of a number of low ecular weight nitrogen compounds. These latter are sometimes called as oup "nitrogen bases" since the nitrogen is present in an amino or no group. The amino acids comprise some of the compounds; and there be very small amounts of urea, creatine, and creatinine, which are untedly intermediates and end products from muscle metabolism. Sevther compounds, also present in small amounts, are anserine, carno-. and carnitine whose structures are shown below. (Carnitine has been wn to have a role in the oxidation of fatty acids.)

O
$$C-OH$$

NH₂CH₂CO-NH-CH-CH₂-C=CH

H N N

Carnosine (β -alanylhistidine)

O $C-OH$

NH₂CH₂CH₂CO-NH-CHCH₂-C=CH

CH₃-N N

CH₃-N N

CH₂CH₂CH₂CO-NH-CHCH₂-C=CH

CH₃-N N

CH₃-N N

CH₃-N N

Water also extracts some proteins and derived proteins from Coagulable protein escapes in the meat juice and during cooking is lated. It forms the precipitate noticeable in the pan juice of roasted The extract also contains gelatin which if not too dilute is not through the increased viscosity or even jellying of the cold Proteoses and peptones are present in small amounts. They originate the hydrolysis of proteins that occurs as the meat ripens. The water B-complex vitamins, particularly abundant in meat, are also extrawater and juice.

Anserine (β -alanyl 1-methyl histidine)

Connective Tissue

This tissue is composed of a few rather large cells scattered thr matrix composed of fibers and amorphous ground substance. The many types of connective tissue; but they all have this common sto of few cells and numerous fibers in ground substance. Connective varies from extremely thin and fragile tissue found between organs to bands and membranes of great strength that form the capsule around organs—the tendons, ligaments, and aponeuroses. A tendon is the band or cord that connects a muscle with some other structure such bone, while a ligament is a band which supports an organ or connection that forms a sheet. Connective tissue also forms the fat or a tissues as well as other specialized tissues such as bone and cartilage.

connective tissue the size, shape, and number of cells varies nously with the source, function, and type of tissue. Unlike most other s, connective tissue has relatively few cells and much intercellular al. Some connective tissues have few cells and these are separated ated. In others they appear more numerous and in some are arranged unite patterns. In all connective tissue the two types of fibers, white zenous and yellow elastic, and the ground substance are of great imnce in determining the characteristics of the tissue.

white collagenous fibers are arranged in bundles of wavy threadlike ents and are coarse and strong but resistant to stretch. In tendons the zenous fibers are arranged in parallel rows with rows of cells between . Since the collagenous fibers are not elastic, tendons also lack elastichey are very strong however.

llow elastic fibers, particularly abundant in ligaments, are fine and hed and distributed in a random pattern to form a network. There w collagenous fibers in ligaments. The tissue is elastic.

e collagenous fibers are composed principally of the protein, collagen, on cooking hydrolyzes to form gelatin. If connective tissue containlarge per cent of collagen is cooked in moist heat for an extended d of time, much hydrolysis occurs, and the tissue almost disappears. clastic fibers, however, are composed of the protein, elastin, which is resistant to moist heat and indeed to most reagents. Cooking has little effect on these fibers.

llagen. In 1952 Bear4 presented a tentative model of the collagen fibril. igure 6.1. Under the microscope collagen fibers can be seen which a diameter of approximatey 100 200 μ (1 μ = .001mm) in tendon and I in the skin. Each fiber is composed of a number of primitive approximately 2 10μ in diameter. By mild mechanical or chemical is the primitive fibers can be divided into fibrils whose diameters are nail that they cannot be discerned with an ordinary microscope but be detected by the electron microscope. Their diameter is measured in of hundreds of Angstrom units (1A = .0001 μ) but even they are far e molecular dimensions. The smallest unit of organization above the cule is the protofibril, a number of which constitute the fibril.

Ita have been accumulating during the past decade with the use of Xiff action methods and electron microscopy for collagen preparations ted to various mild conditions. These data give some indications of ape of the collagen molecule and the manner in which molecules are 17ed in the protofibril. Bear has suggested a tentative model which

re briefly described.

e fibril appears to be a very long thin body containing bands and

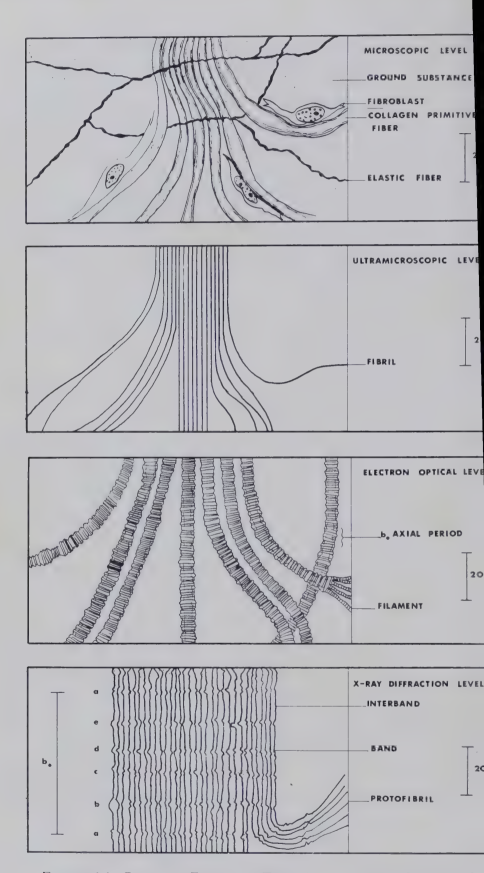
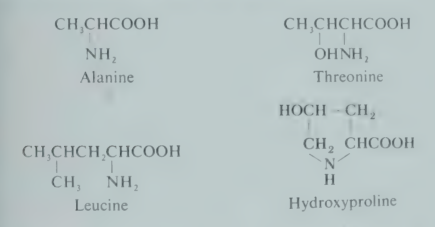


Figure 6.1. Collagen Fibers and Their Arrangement. Courtesy of R. Bear.

sands at definite distances. This series of bands reappears at 640Å at the length of the fibril and appears to have significance as far as the reement of the side chains on the collagen molecule is concerned.

e collagen molecule is a polypeptide chain with a very high percentage veine residues. Beef collagen, which has been studied most extensively. ...ins 19.9 per cent glycine. It contains a very high (64 per cent) perage of amino acid residues which have nonpolar side chains (glycine, me, alanine, the leucines, valine, phenylalanine, and methionine). ther or not these amino acid residues are arranged in definite patterns not been determined, but Bear believes that they are arranged in a mte pattern as far as type, with a section of nonpolar side chains wed by polar ones capable of hydrogen bonding and salt links. Polar chains are those containing hydroxyl groups (hydroxyproline, onine, serine, and tyrosine), those containing acidic groups (glutamic and aspartic acid or their amides), and those containing basic groups ne, arginine, histidine, and hydroxy-lysine). Hydroxyl and acidic rs are active in hydrogen bonding, while acidic and basic groups in e approximation form salt bridges. Collagens are unusual in their poson of fair amounts (from 5 to 13 per cent) of hydroxyproline.



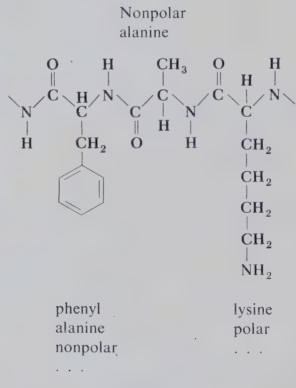
ne collagen molecule is coiled and the protofibril is made of parallel cules coiled or twisted together to form a helix. The protofibril is red of many molecules in length; and although the manner in which the of one molecule abuts the end of another is unknown. Bear believes the molecules do not overlap in the protofibril as they do in some there but instead lie parallel to one another along the length of the eptide chain.

to bands and interbands are believed to be caused by the side chains of icent molecules and the way in which they mesh or crowd one her Bear suggests that the interbands are regions in which the rel-

atively small nonpolar groups can mesh with one another in such that the polypeptide chains are able to lie quite straight in the linterbands are areas where the relatively large polar side chains and react with one another, producing a kinking of the polypeptid A short portion of the coil, straightened out, is shown in the il (Figure 6.1).

Collagens from animals in widely separated phyla show many ties, but some differences. Collagens from mammals have bee most extensively with fishes next; a few examples from other pl been examined. The preliminary work indicates that the collag closely related animals have marked resemblances, while the fart they are in the animal kingdom, the greater are the differences in The gross structure of the molecules of animals from different pears to be similar, but the amino acid content is different.

Elastin. Elastin is the protein that forms the yellow elastic fiber collagen, it is not hydrolyzed on boiling with water and consectissue that contains a considerable number of yellow elastic fiber little softening or dissolving upon cooking. These fibers are abutendons and ligaments which are often trimmed away before cooked. The human Achilles tendon contains 20 times more elastic fiber.



Polypeptide Chain

gen and in the ligamentum nuchae, which extends down the neck of and helps hold up the head, there is approximately 5 times as There has been some study of the structure of elastins by the ods applied to collagens. They appear to be linear proteins but of erent type than the collagens. Beef elastin differs from beef collagen amino acid composition. It is unusually rich in amino acids with olar side chains (glycine, the leucines, valine, phenylalanine, and prowith 78 per cent of the molecule made up of these amino acids. It is ar to collagen in its small content of histidine, cystine, tyrosine, and tophan.

ne ground substance of connective tissue varies from a soft jellylike s to a tough matrix. In cartilage and bone the ground substance is quite rent from that in other connective tissue. It is principally composed of er in which salts, glucose and other molecules are dissolved and which eves its jellylike characteristics from the presence of heteropolysacnces, colloidally dispersed. In cartilage, chondroitin sulfates are most ant while in soft connective tissue and skin, hyaluronic acid occurs.

p. 80.)

dipose Tissue. This is a specialized type of connective tissue in which e is rather than the intercellular material are most abundant. Fat cells ound scattered through, or in groups in loose connective tissue, but the "adipose tissue" is reserved for those tissues containing large deposits t. Adipose tissue varies in color from very light cream to dark yellow. o'ten classed as white and brown adipose tissue. It occurs in the subneous connective tissue, in the region of the kidney, in the omentum, around and between some of the muscles and organs.

at a fat cell is filled with a large single vacuole of fat, and the cytom and the nucleus are pressed by this vacuole close to the cell wall. fit cells are arranged in groups or lobules with delicate connective e that is rich in fibers and many blood capillaries separating the

dipose tissue is by no means composed entirely of fat. It is true that bef lipid is neutral fat, but the cells possess proteins of many kinds. r, salts, and other compounds commonly present in cells in small ants. The neutral fat present in the adipose tissue is characteristic of p cies; the assortment of fatty acids follows fairly closely a species 3ut in the carcass some variation does occur in the composition of ifferent tissues. In general, the outer layers of adipose tissue confa' with higher iodine numbers and lower melting points than the inner 's I pid occurs in the vacuoles in the liquid state, and the fat near the is at a slightly lower temperature than that in the interior of the living

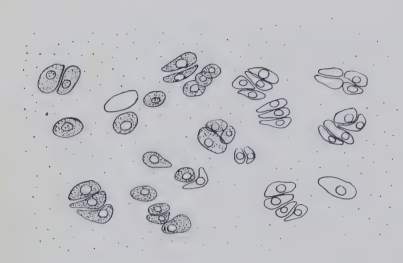


FIGURE 6.2. SRIB CARTILAGE Matrix is slight × 240. (Redraw Sharpey-Schafer H. M. and Short "Schafer's Ess Histology," Green and Co. 1956.

body. It has been suggested that in the living animal the fatty acid of the fat deposited is as saturated as it is possible for it to be and main in the liquid state. Some observations on the effect of extern perature appear to support this. The higher the skin temperation higher is the melting point of the fat deposited.

It is well known that the properties of lard produced from differences of a hog vary and this fact is utilized in commercial fat produces p. 52 for a discussion of the production of fat from animal sources.

Cartilage. This is also a specialized type of connective tissue; other connective tissue cartilage (Figure 6.2) is composed of cells and ground substance. The cells occur for the most part in island lacunae, surrounded by ground substance in which the fibers, be lagenous and elastic, are embedded. The ground substance, is a gposed of chondroitin sulfate, chondromucoid, and albumoid in water

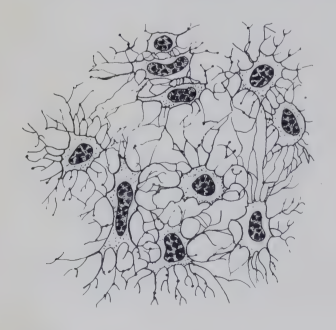


FIGURE 6.3. BONE CEL THEIR CANALICULI IN TH BRANE BONE OF MOUSI Maximow and Bloom) × 10 tesy of W. B. Saunders Co.

TABLE 6.1. COMPOSITION OF BONES*

	Cattle bones
CaCO ₃	7.07
$Mg_3(PO_4)_2$	2.09
$Ca_3(PO_4)_2$	58.30
CaF ₂	1.96
Organic	30.58
Total	100.00

a mted from Edelmann, R. H., Mohler, J. R. and Eickhorn, A., "Textbook of Meat Hygiche," Leaeer, Philadelphia, Pa., 1943.

Some. Bone (Figure 6.3) is also a specialized connective tissue. Like tilage, it is composed of cells located in lacunae, fibers, and ground estance in which tiny crystals of salts are deposited. In bone usually only cell occurs in each lacuna, and it can communicate with others through cries of small canaliculi which traverse the matrix. The fibers are numerand are collagenous. They are arranged in small bundles and run in defee patterns. Bones are traversed by numerous canals, some of which run pendicular to the shaft (Haversian) and some of which run obliquely or the angles (Volkman's). These canals make the bone porous and less use. The ground substance is composed of the proteins ossomucoid and calbuminoid surrounding small crystals of salts. The most abundant is calcium phosphate, but calcium carbonate, magnesium phosphate, calcium fluoride are also present. See Table 6.1.

he proteins in bone are for the most part unidentified. Food gelatin is nufactured from fresh bones by boiling them in water, a process conting collagen to gelatins. In some processes the bones are boiled in very ate hydrochloric acid, the phosphate is precipitated and the acid tralized with lime water, after which the product is dried. The product alled "ossien" and on extraction with hot water yields gelatin. The "is "ossomucoid" and "ossoalbuminoid" have been assigned to the prosin the living bone, but little is known about them.

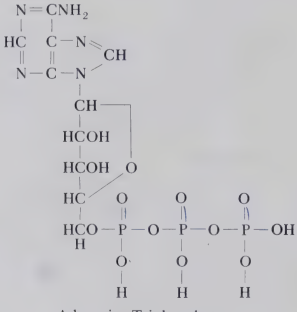
STMORTEM CHANGES

when an animal dies, the skeletal muscles stiffen in rigor mortis and remains this condition for a period after which they soften and become sle again. The speed with which rigor develops and the length of time exists is variable. It is widely believed that the onset of rigor is exists is variable. It is widely believed that the onset of rigor is exists is variable. It is widely believed that the onset of rigor is exists is variable. It is widely believed that the onset of rigor is exist in the temperatures and delayed by low ones. But Bate-Smith and that the temperature of the carcass was of little importance computed to other factors and Stewart, Lowe, Harrison, and McKeegan* concept to other factors and Stewart, Lowe, Harrison, and McKeegan* concept to other factors and Stewart, Lowe, Harrison, and McKeegan* concept to other factors and Stewart, Lowe, Harrison, and McKeegan* concept to other factors and Stewart, Lowe, Harrison, and McKeegan* concept to other factors and Stewart, Lowe, Harrison, and McKeegan* concept to other factors and Stewart, Lowe, Harrison, and McKeegan* concept to other factors and Stewart, Lowe, Harrison, and McKeegan* concept to other factors and Stewart to ot

firmed this for chickens. They found the onset of rigor was graduchickens studied and usually varied between 1 and 2 hours. Howe chicken developed rigor mortis in 5 minutes. Rigor mortis is important products since muscles cooked while still in rigor are much than if it is allowed to pass before cooking.

The stiffness that develops when muscles pass into rigor is the rechanges in the proteins. Living muscle fibers contain protein in pliable gel. During rigor this gel stiffens, but when rigor passes, the again becomes soft and pliable. Finally during cooking, another called *rigor caloris* occurs and the proteins stiffen again. The in the proteins of muscle are still incompletely understood although ber of theories to explain rigor have developed. While some knowle accumulated, there is still much ignorance concerning the chemical cinvolved.

In a living muscle changes occur in the proteins actin and when a muscle contracts. Szent-Györgyi⁴⁹ has presented a rathe plex theory to explain the many reactions that lead to a change i and myosin, the formation of a complex, actomyosin, and the cont of these molecules. It would take too much space to give the theor and it is constantly changing; but the fact that investigators have so far in their study of muscles that the formation of a theory is p is encouraging. The understanding of the chemistry of the muscle and after death of the animal will not be far behind. The energy for traction in living muscle is supplied by adenosine triphosphate,



Adenosine Triphosphate •

on possesses a high energy phosphate bond in the second and third

sphate.

pe energy for contraction and its accompanying heat production comes apally from these high energy phosphate bonds. Resynthesis of AIP in tying cell is at the expense of glycogen which passes through a longes of enzymatic reactions to form pyruvic acid. The pyruvic acid is then ized by way of the citric acid cycle to carbon dioxide and water, and it bursts of energy are released as each carbon and hydrogen are oxid. This series of reactions is completely discussed in any modern texts on biochemistry. The research that has led to the elucidation of this es of reactions is a triumph for modern biochemistry. The over-all series is following:

$$2(C_6H_{11}O_5)_n + 5[O] \longrightarrow 4n CH_3COCOOH + H_2O$$
Glycogen Pyruvic Acid
$$CH_3COCOOH + 5[O] \longrightarrow 3 CO_2 + 2 H_2O$$

the transformations of glycogen to pyruvic acid require a small amount oxygen, but the citric acid cycle for the oxidation of pyruvic acid to con dioxide and water requires more (5 atoms of O for each COCOOH). The oxygen is carried to the muscle by the blood. When itemand for oxygen in a contracting muscle exceeds the supply, a contractled "oxygen debt" develops and some pyruvic acid is reduced to cacid to supply oxygen.

or an animal is killed and circulation of blood ceases, the degradacy glycogen continues, small amounts of intermediates accumulate, with the influx of oxygen stopped, lactic acid is the most abundant uct. Muscle contains some buffers that neutralize the first lactic acid red. Later as more and more forms, the pH of the tissues begins to fall. amount of glycogen stored in the muscle at the moment of death conthe amount of lactic acid formed and, consequently, the ultimate pH meat. A well-nourished muscle will have as much as 1 per cent glycotom 1.1 per cent lactic acid, and drop to a pH of 5.6. The pH never below 5.3 because some enzymes become inactive at this low pH.

tors that influence the amount of glycogen in an animal have been the at of a number of studies since a low pH, we shall soon see, is usually able in meat from many standpoints. Whether or not animals are fed 4 hours before slaughter affects the amount of muscle glycogen and the resultant pH of the meat. It is only in the well-fed animal that

muscle cells contain maximum glycogen. The amount of exercise an has just before slaughter is important. An animal that is chased or or driven into the killing pen, will use up much of the muscle glycoging the vigorous exercise, and without a rest period to allow for r the supply, will produce meat in which the development of a low pl possible.

The series of chemical reactions occurring after slaughter p enough heat to cause a rise in the temperature of the meat. The body temperature is 99.7°F in cattle, but shortly after death, the temperature of a round of beef may rise to 103°F. The fresh meavery slowly even in a refrigerator because of the continuing product heat. This heat production, called *animal heat*, has been recognized beers and butchers for hundreds of years. It has been often consomething mysterious and mystical. Farmers will often say that me not be eaten until the animal heat has escaped, that it is not fitting meat with animal heat, or make similar statements. Because of the ration of animal heat, freshly killed meat is seldom eaten. A short perhanging during which rigor develops and passes and during which ripening may even begin improves the meat before cooking.

Both creatine phosphate and adenosine triphosphate are hydrolyging rigor begins to develop. Creatine phosphate forms creatine and phosphate of the muscle enzymes, a phosphatase capable of hydrolyzing sine triphosphate to inorganic phosphate and adenosine diphosphate comes more active as the pH drops to 6.5 and below. The adenosphosphate is then further decomposed to ribose, phosphate, ammor hypoxanthine (from the adenine). Contraction is no longer possible muscle, and a tension develops which is incapable of relaxation rigor passes.

Both Bate-Smith³ and Szent-Györgyi⁴⁹ have proposed theories of mortis that include reactions of the proteins of muscle and chathem. Bate-Smith believes that the disappearance of adenosine to phate is of great importance in the development of the stiffening of mortis. Szent-Györgyi points out that creatine phosphate recently his shown essential for relaxation of contracted muscle, and he believes appearance may be of prime importance in the stiffening of rigor more contracted muscle.

The sarcolemma, the membrane around the muscle cell, shows a dat this time in electrical resistance, and a free diffusion of ions. The change in the semipermeable membrane indicates that not only protein of the muscle fiber changing but the substances that make membrane, the sarcolemma, might be changing chemically. The electronic of the muscle as a whole changes also.

he ultimate pH attained by the meat has important effects on other perties of the meat. Meat with a high pH is darker in color than normal, ny to touch, and does not allow salt of curing pickle to penetrate read-It is also difficult to express juice from meat with a high pH. The dark that sometimes develops in beef carcasses has been studied by several ups because it is of considerable economic importance. Consumers are ctant to buy beef that is darker than normal. The pigment of muscle, valobin (p. 189), is present in the same concentration in dark cutting as in normal; but the light passes into and is reflected from deeper rs of the muscle, giving a darker appearance. A second factor appears be a difference in the openness (or closeness) of the grain of the meat. high pH the swelling of the protein is greater, and because of this the ity of oxygen to diffuse through the tissue is probably decreased. Less gen causes a decrease in the proportion of oxymyoglobin to myoglobin, a darker color. It has also been shown that the consumption of oxygen dark meat is higher than by light meat, but the significance of this obation is unknown.

Meat that develops a relatively high pH is difficult to cure properly bese the pickling salts do not penetrate the meat at a normal rate. Callow studied the relation of the pH of meat to its electrical resistance and nd that between pH 5.8 and 6.0 there is a rapid increase in resistance. It is also the range of pH's in which the grain of the meat shifts from n to closed. These observations have been interpreted to mean that ling of the protein occurs in this range and that the passage of ions is impeded. It is thought that the increased resistance to the passage of electric current reflects this difficulty of ion migration.

ratts⁵⁴ points out, however, that a relatively high pH is desirable in at which is to be frozen since in this range of pH's discoloration and rate of fat oxidation are inhibited and the drip losses are minimized.

allow has also related the pH of the meat to the rate of growth of teria in Wiltshire cured hams. He showed that the occurrence of taint in hams was correlated with the electric resistance and with the pH. Hams flow pH showed lower electrical resistance and a greater freedom from the concluded that with lower pH the penetration of the pickle was a rapid and that the bacteria were not able to grow as readily in the excid medium. Ingram showed that this was true by isolating organ-from spoiled hams and measuring the bacterial growth at various Growth was very low at pH 5, increased rapidly with rising pH, and

hed a peak at near pH 8.*

Ripening or Aging

After the passing of rigor mortis, meat becomes progressive tender, juicier, and more flavorful. The speed with which this rip aging occurs depends on the time the carcass is kept and the temp Changes occur quite rapidly at room temperature but more slow frigerator temperatures. Only well-finished carcasses with a good fat are satisfactory candidates for ripening since a growth of mol must be trimmed away before the meat is cooked appears on the Poorly finished carcasses rapidly develop off-flavors and putrify. discusses the work which Hoagland, et al. carried out many years the ripening of beef stored just above the freezing point for 17 to 17 The organoleptic qualities of the meat were judged by a panel whi sidered the meat stored 15 to 30 days better in tenderness and flav the fresh meat. After 45 days the meat began to develop off or gar vors. A taste for well-ripened meat varies with different groups of Well ripened meat is preferred for the most part by the English but Americans.

Some aging occurs during the time which must inevitably elenormal commerce between slaughter and preparation of the meat table. Purposeful aging is costly because of the use of expensive retor space, the additional trimming required to remove mold, a shrinkage caused by evaporation. It is used to a limited extent aronly for prime and choice grades of meat.

A recent study⁴⁴ shows that aging beef at elevated temperature high humidity, with air velocity 5 to 20 lineal ft per min and with violet radiation to control microorganisms for 2 or 3 days produce equal in quality to that aged in a refrigerator 12 to 14 tlays. Quere judged on tenderness, flavor, aroma, and juiciness.

Although a number of studies of composition before and after have been completed, the numerous changes that occur on ripeniaging still are not fully known. The reactions are the result of an action in the cells—reactions that fragment large molecules through action of enzymes and reactions that result from changes in cell conceparticularly the change in pH. Ginger, et al. 18 determined the conted distribution of nitrogen compounds and particularly the amino arginine, lysine, leucine, tyrosine, histidine, and glutamic acid in recooked beef and in the drippings from the meat before and after agin cut used was rib steak. Aging caused an increase in the free amin nitrogen. The nitrogen in drippings of freshly slaughtered meat was mostly as nonprotein nitrogen (NPN) with a large proportion of amino nitrogen. After the beef had aged for two weeks, drippings of

nerease in the amount of NPN. The amino acids histidine, leucine, sine, glutamic acid, and lysine were present in a bound form. Histidine ounted for as much as 26 per cent of the NPN in the extracts. It was luded that part of the bound histidine is present as carnosine.

olov'ev and Piulskaya46 studied the change in concentration of a few ponents of beef during ripening. In the first study the flesh of cows was ed at 0°C to 40°C for 144 hours and samples were taken and anad at intervals. Actomyosin showed a drop in solubility during the first nours and then a slow rise. The abstract reports that the "activity" of actomyosin increased from 6 per cent at 12 hours to 56 per cent at 48 rs and that there was some increase during resolution of rigor mortis. P is quickly destroyed and they found that at 12 hours 90 per cent of ATP was gone. The first report likewise confirmed the fall in free glycoand reported that there was also a decrease in "difficultly extractable ogen" (quoted from abstract). In the second report45 beef stored at . 8° 10° C, and 17° C was analyzed daily for some constituents as well rganoleptically. Optimum changes occurred in 10 days at 0°C, in 4 days 10°C, and in 3 days at 17°C. The chemical analysis paralleled these anoleptic changes with maximum increases in volatile fatty acids, volareducing substances, total nucleotide N, adenylic acid, free purine, and oxanthine. These observations serve to confirm in part the hazy idea ripening is a period when autolytic changes occur and large molecules tinue to form small ones although syntheses of the large ones that ntain the tissue during life have stopped.

tinues even in frozen meat. Sizov measured the myosin and its enzyme vity for adenosine triphosphate in rabbit meat stored at 14°C. He that at the end of 5 days the amount of myosin had fallen to 58.5 cent and eventually it disappeared. At the end of 22 days the enzyme vity was only 30.3 per cent of the fresh meat. Drozdov studied beef er a number of storage conditions and found evidence of autolysis in all plus Some were stored from 2 to 4 months at 10°C and showed o vsis, accumulation of lactic acid, and decomposition of phosphates.

Harrison, et al.²² reported studies of four muscles removed from steers 14 hours after slaughter. The aroma and flavor reached a maxi in 10 days and decreased after 30. Tenderness as judged by the shear and the taste panel increased particularly during the first 10 days. Acrose during the first 2 hours.

Paul, et al.³⁹ compared tenderness in beef steaks and roasts held varying lengths of time in cold storage. They used the semitendinosis the biceps femoris of 2 prime, 2 good, and 2 commercial carcasses, meat was hung 0, 5, 12, 24, 48 to 53, and 144 to 149 hours. Althouse steaks showed decreased tenderness up to 24 hours, with longer tenderness increased; whereas roasts were least tender at 0 time and sho increasing tenderness on aging. The tissues were studied histologically showed evidence of the occurrence of rigor in the roasts cooked at 0 the As rigor passed, breaks and areas of granulation occurred in the mufibers.

COLOR OF MEAT

The principal pigment present in muscle cells is myoglobin, a red cogated protein closely related to hemoglobin of the red blood cells. We meat is prepared for market, it is not completely bled and some red blood cells.



FIGURE 6.4. MODEL OF M GLOBIN MOLECULE. Construater mapping the globular recule with X-ray diffraction stu Polypeptide chains are white gray patch is the heme. Courte Dr. J. C. Kendrew,

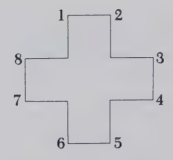
remain in numerous blood vessels which traverse the muscle tissue, se contain hemoglobin. Small amounts of colored enzymes which are repigments also occur in muscle cells. They are cytochromes and raidases. However, most of the color of meat is caused by the pigment remuscle cells, myoglobin.

tyoglobin has been prepared from beef and pork, as well as other cles, but it has not been studied nearly as extensively as hemoglobin. It is a globular protein composed of one heme moiety for each ecule of protein while hemoglobin contains four heme moieties and of protein. The molecular weight of myoglobin is just one fourth that emoglobin, 17,000 versus 68,000. Likewise myoglobin shows a differing amino acid composition and solubility. It can be saturated with gen at lower oxygen pressures than hemoglobin and its reaction with non monoxide and nitric oxide and the stability of the products are also inct. The porphyrin, heme, in each molecule is the same, but the prosare different and doubtless the manner in which they are combined is trent. Mapping by X-ray²⁹ analysis suggests a structure for myoglobin wn in the picture. (Figure 6.4).

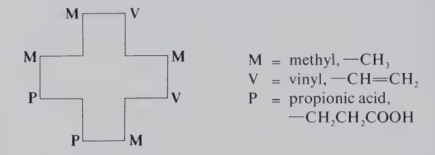
he porphyrins are a group of compounds that form the prosthetic ups of many colored conjugated proteins. In chlorophylls (Chapter 7) porphyrins contain magnesium, while in some forms of life such as lusks the porphyrins contain copper. In hemoglobin and myoglobin metal held by the porphyrin is iron. The porphyrins are ring comnds composed of four pyrrole rings held together by CH or methene ups. A convenient shorthand formula for the basic porphyrin structs shown below in which each of the corners is occupied by a carbon tCH.

Porphyrin Nucleus

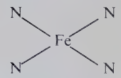
ther abbreviation of the formula occurs when it is written as a cross. phyrins differ from one another in the side chains commonly attached he "corners of the cross."



In hemoglobin and myoglobin the porphyrin "corners" hold not vinyl, and propionic acid.



It will be noticed that in the structure which shows the nitrogens, to the nitrogen atoms have three bonds and two have only two bonds third valence of each of these latter nitrogen atoms is satisfied by bination with ferrous iron. The iron atom has the ability to coordinate share electrons from nitrogen atoms, and in the hemoglobin molecular may be thought of as being attached through an ordinary covalent both two of the nitrogen atoms and through a coordinate covalent bond to other two. The whole structure is in resonance, and it does not make the which of the nitrogens is considered to be held by ordinary valence which by coordination since in any case the bond consists of a parelectrons.



Iron has a coordination number of six.

Heme, ferroprotoporphyrin, of hemoglobin and myoglobin can bine with a number of organic nitrogen compounds. Among then histidine, although it does not react with arginine and lysine. In h globin a nitrogen (one of those circled) from the imidazole ring of hist is believed to coordinate with the ferrous iron and hold the heme in

$$CH_{3}C = CCH = CH_{2}$$

$$HC-C = CH$$

$$CH_{3}C = CH$$

$$C = CH$$

$$C = CCH_{3}$$

$$N-Fe-N$$

$$HOOCCH_{2}CH_{2} = CCH - CH_{2}$$

$$HOOCCH_{2}CH_{2} = CCH_{3}$$

$$Heme$$

$$Ferroprotoporphyrin$$

e protein molecule. The sixth pair of electrons comes from the oxygen nolecule of water in hemoglobin or a molecule of oxygen in oxyhemon.

Histidine

our nitrogens of the porphyrin are in one plane with a nitrogen of nidazole ring above the iron, and H₂O or O₂ below. There is likewise sibility that the carboxyl group of the propionic acid side chains may ked with protein, but this has not been proved.

Imidazole

unique property of hemoglobin and myoglobin is the capacity to m lecule of oxygen without a change in the oxidation of the iron. Button is readily reversible and is most important in the transfer gen to the tissues in the living animal. In meat it accounts for the n color that occurs when meat is cut. The myoglobin-oxymyo-

globin shift occurs readily when myoglobin is exposed to oxygen. The change in color occurs when meat is cut. Oxyhemoglobin and ox globin have a very bright red color, while hemoglobin and myoglob more purplish red. The bluish color of meat is caused by the large am of myoglobin (and some hemoglobin) present, but when the meat is cut the surface exposed to air, the color becomes bright red as oxymyog is formed. Other heme pigments of muscle, peroxidases and cytochr cannot combine in this fashion with oxygen.

When hemoglobin is treated with a strong acid or hydroxide, the p and the porphyrin are separated.

Hemoglobin + HCl → Globin·HCl + Ferroprotoporphyrin (heme)

The ferroprotoporphyrin, heme, is rapidly oxidized under most cond to ferriprotoporphyrin, hemin, in which the iron exists in the ferric The crystallization of hemin is a common and delicate test for the dete of blood. Hill and Holden demonstrated that the globin formed by th action will dissolve in water near pH 7 and recombine with heme, protoporphyrin, to form hemoglobin, or with hemin, ferriprotoporph to form methemoglobin.

Ferroprotoporphyrin (Heme) + Globin → Hemoglobin
Ferriprotoporphyrin (Hemin) + Globin → Methemoglobin

If hemoglobin is treated with oxidizing agents, it is changed direct brown methemoglobin.

Hemoglobin oxid, Methemoglobin

Methemoglobin can be separated into ferriprotoporphyrin (hemin) globin.

Interestingly enough the globin from different animals will combine any heme, no matter what its source. The hemoglobin produced by combination always resembles the hemoglobin of the species from very the globin was prepared. It is the protein portion of hemoglobins different animals that is species specific. Edmundson and Hirs¹² have slethat sperm whale myoglobin contains the following proportion of a acids: $Glu_{19}Asp_8Gly_{11}Ala_{17}Val_8Leu_{18}Ileu_9Ser_6Thr_5Met_2Pro_4Phe_6Tyr_9His_{12}Lys_{19-20}Arg_4$.

Methemoglobin is formed by the reaction of a number of oxid agents such as peroxides, ferricyanides, and quinones with hemoglobin has a brown color, as does metmyoglobin. Metmyoglobin is simformed by oxidation of myoglobin and is consequently believed to be

ogue of methemoglobin. In both of these conjugated proteins the iron in the ferric state. Methemoglobin is normally produced in small ants in the living animal, but it is again reduced to hemoglobin. The ion has been studied because it is important in some pathological rions and in the preservation of whole blood. In meat the formation ownish and gray pigments, or the dulling of the bright red of fresh as it ages, may be caused by metmyoglobin formation.

owning of meat may sometimes occur through oxidation of heme after ciation from the protein. Ferroprotoporphyrin, heme, is much more by oxidized than hemoglobin and probably more rapidly than myoglo-Often when meat darkens or browns, conditions are right for the turation of the protein and the formation of free ferroprotoporphyrin, Unless conditions are such that oxygen is excluded, and this usually true for meat, the ferroprotoporphyrin is quickly oxidized to ferriporphyrin, hematin. When cooking occurs, heat coagulates or detes the protein; when meat is marinated in acid such as vinegar, detation occurs. Some agents allow the heme to become disengaged the protein and consequently be more susceptible to ferric oxidation. Ting, salt, acid, or ultraviolet light are capable of denaturing protein are known to affect the color of meat.

casionally after meat has been cut for some time, the color of the ce will deepen but it will remain bright red. In a study of packaging, rock and Wallace³⁰ found that this deepening is caused by dehydratif the surface of the meat and a consequent concentration of the ent.

een pigments occasionally occur on meats. The green pigments ed from the porphyrins that have been studied are all products in the alpha methene carbon bridge is attacked. In the living animal protoporphyrin, heme, is normally changed into green pigments and me animals they are discharged in the bile as the bile pigments. In animals green pigments are changed to red and then excreted in the Choleglobin is a compound in which the porphyrin is still conjugated protein (and in which the iron remains as ferrous iron) but where pha methene bridge is partially oxidized but not broken. Verdoheme compound formed when oxidation of the alpha methene bridge is sive enough to break it. The green pigments formed in meat may be reompounds.

ent situations. Jensen²⁷ has shown that the hydrogen sulfide produced rtain bacteria in the presence of oxygen can add directly to the porning and form an analogue of choleglobin. Hydrogen peroxide-

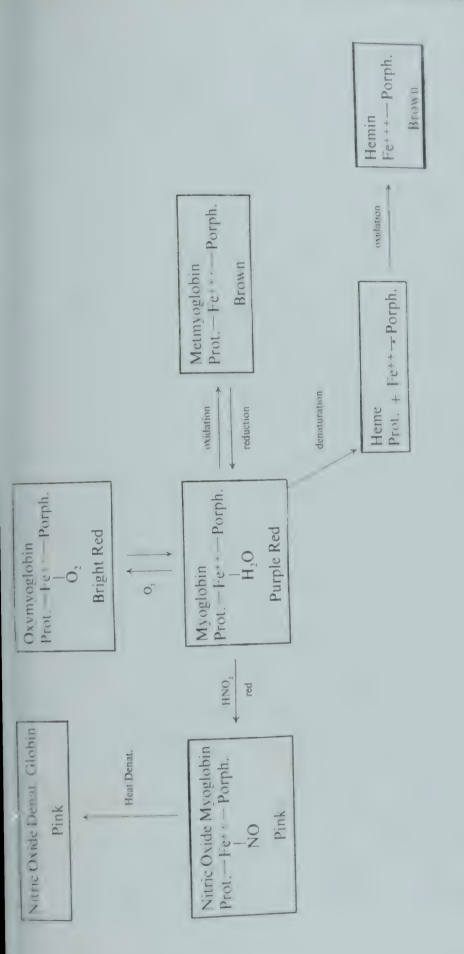
forming bacteria will bring about the same type of reaction with mation of choleglobin itself. In fresh meat any hydrogen peroxide is rapidly destroyed by the action of the enzyme catalase but i meats, catalase is absent and greening of the meat occurs readily compounds are able to bring about similar reactions. The one whosen most extensively studied is ascorbic acid which, of course, is n present in fresh meat at low concentrations.

Both Jensen²⁶ and Urbain⁵¹ say that the identity of green pign unknown. In 1936 they published work on the spectra of pigments of on exposure of nitric oxide hemoglobin to bacteria. Five different for green derivatives were obtained. They have shown that numero pathogenic organisms can produce either oxidizing agents or hydrofide and that either type of product will react with nitric oxide hem to form green pigments.²⁷

Iridescence or mother of pearl effect is sometimes observed in cured or cooked meat. The play of color changes as the point of the source of light changes. Iridescence is commonly produced it soap bubble films, or when white light passes through some sort fraction grating and is broken up into its components to form ra In the case of iridescence of meat it is believed that light is broke this manner as it passes through a film of fat on the surface fibers fat is removed, the iridescence disappears; if it is again added, the colors reappears.

The color of meat is also directly related to the development of rain meat. It has been found that myoglobin and hemoglobin acceleronset of rancidity in the fat of meat. As oxidation of the fatty acceeds there is also an oxidation of the myoglobin to metmyoglobin.

In the cured meat nitric oxide hemoglobin and nitric oxide my are formed. Nitric oxide hemoglobin has been studied and found compound very similar in structure to oxyhemoglobin. The nitric



*From Watts, B. M., "Oxidative Rancidity and Discoloration in Meats," Advances in Food Research, 5, 1-42 (1954).

NO, is held by coordinate valences by the ferrous iron in the same was a molecule of oxygen is held in oxyhemoglobin. The two compound very similar bright red colors. When cured meat is cooked, the pink persists because nitric oxide hemoglobin (or myoglobin) is denatured nitric oxide hemochromogen which is also pink.³

CURED AND SMOKED MEATS

Long ago when man began to cook his food and to live in some k shelter, he soon learned methods of storing meat. Probably he firs covered the method of smoking meat by hanging it near the roof cave or tent where it slowly dried and became smoked. Later he that the addition of salt to the meat (called pickling) prevented its faction and salt became one of man's most valued possessions. Salt r developed and a man's wealth could be measured by his supply of salt sen tells us that in Homer's time (about 1000 B.C.) both smoked cured meats were generally consumed. Today with adequate refriger and rapid distribution, there is little need for curing and smoking preserving measures. But our appetites have been developed for the ticular flavors which smoking and curing develop, and these me are still widely practiced. In early times spices were often added ticularly to the comminuted meats, to disguise the flavor of "high slightly putrid meats. Today we relish these spices even though the must always be fresh and clean. During the middle ages in Europe small region developed its own particular type of sausage. Here in Am our cuisine is the result of food customs from all of Europe and to a ited extent from the rest of the world. The number of sausages on the sumer market today is therefore enormous, each with a subtle differen its recipe. From a chemist's standpoint, they are all very similar.

Since the need for preservation of meat by curing and smoking is la eliminated by our modern methods of refrigeration, the sanitary condition of the meat packers, and the rapidity with which distribution occurs, no longer necessary to use as much salt in the pickle. Gradually the erence of American consumers has developed for mild cures—one which the salt content is kept very low. Meats that are mild-cured are protected sufficiently by the salt to be stored at room temperature, by ways require refrigeration.

There are four general methods of curing meat: (1) dry cure, (2) pcure, (3) injection, and (4) comminution and mixing. The Meat Insper Division, FDA, allows five substances for curing: sodium chloride, so nitrate, sodium nitrite, sugar, and vinegar. In the dry cure a mixture

first four compounds is rubbed on a cut of meat, and it is held under geration for some days while the compounds penetrate the tissue. In pickle method the compounds are dissolved in water, and the cuts of it are submerged in the pickle for a period of time. Here the water nod is now widely used in the United states, often in combination with of the other methods. All meat contains arteries and veins and their sches—a complete vascular system. Pickle can be pumped in through of these vessels and will be carried through the meat. In curing ham common practice to pump pickle in through the main artery (the iliac) ne leg. The needle is also inserted into the muscle and pickle forced the muscular bed. In the fourth method the meat is simply chopped and the curing compounds mixed with it. Cure is rapidly effected bese contact is immediate. Vinegar is not used in commercial pickle.

he term "corn" is an Anglo-Saxon word for a small grain, often apby the people to the common cereal grain of the area—for example, ne United States corn is maize, in England, wheat. Since in early efto preserve meat the salt used was granular, the pickling or curing ess came to be called "corning." The meat so cured is sometimes

:d "corned."

ne action of salt, NaCl, in preserving meat is to inhibit the growth icteria. Jensen²⁶ reviews some of the rather sizeable body of research covers the effect of salt on microorganisms. Salt shows a selective on and does not inhibit the growth of all microorganisms. Many yeasts, xample, grow in media containing as much as 15 per cent salt. It has demonstrated that the effect of salt depends not only on its concentrabut on the presence of other compounds in a nutrient medium. Some nisms will not grow in the pickle but will grow on the surface of the covered by the pickle since the nutritional environment is different . Salt also has a detrimental effect on meat. It is well established that accelerates oxidative rancidity. When a dilute salt solution is brought ontact with fat, it may actually retard oxidation; but if the solution is ally evaporated so that a film of salt is deposited on the fat, rancidity eatly accelerated. Gaddish has shown that in bacon the greater the entration of salt, the more rapid the development of rancidity. In un-I meat, salt also accelerates the formation of brown methemoglobin; red meat, when nitrite is added, the opposite effect is noticed. avors the development of the bright red color. The nature of the efs not known but it may well be that it is through the denaturation e globin. Salt is a denaturation agent.

re use of nitrates in curing meats originated from the discovery dur-

ing the late Middle Ages that the color of meat becomes fixed when is added. The nitrate is reduced to nitrite by nonpathogenic organ the pickle. At the pH of the meat, 5.4 to 6.0, nitrite exists as nitrou HNO₂. There are also reducing conditions in the meat because of the ence of readily oxidized compounds formed under conditions of debt following slaughter. The nitrous acid is reduced to nitric oxid which then reacts with myoglobin to form nitric oxide myoglobin.

- (1) $NO_3^- \rightarrow NO_2$ (bacterial action)
- (2) $NO_2^- + H^+ \rightarrow HNO_2$ (at pH of meat)
- (3) HNO₂ → NO (reduction by compounds in meat)
- (4) NO + myoglobin → NO(myoglobin bright red)

Saltpeter is the old name for potassium nitrate and Chile saltpe sodium nitrate. In meat packing houses either one is commonly "saltpeter" although since the sodium salt is cheaper, it is usually The nitrate also has some bacteriostatic effect.

Watts⁵⁴ points out that the series of reactions is not quite as simmany cases as that presented above. When nitrite comes in contact oxyhemoglobin, the first reaction is the formation of methemoglob trate, and oxygen. Thus when ground pork sausage is mixed with salts, the color immediately turns gray as metmyoglobin is formed reaction is very rapid at the low pH of meat. When the meat is haduring smoking, the red color of the nitric oxide hemochromogen dev Watts also believes that there is a good possibility that other intermed are involved and that the reaction on heating is not as simple as the given below.

NO-myoglobin heat NO-hemochromogen

The development of the proper color during curing depends on the mation of nitrite, and this in turn depends on the presence of or groups of bacteria which slowly reduce the nitrate to nitrite. When recognized that nitrite was the color-fixing agent, it became obvious the problem with undercuring—poor color development—could be so by the addition of nitrite as such to the pickle. In 1927 the use of so nitrite was allowed. Today the common practice is to use both nitrate nitrite since nitrite alone often does not give good color. If nitritused alone, the concentration of nitrous acid is so great that it decommon and N_2O_3 is lost from the pickle. The action is not prolonged of tained during the whole of the time of pickling, so that a combinate both nitrate and nitrite is, in general, more successful.

duces conditions during curing and storage that insure the best color conserve protein. Ordinarily, sucrose is added to the pickle. In the ence of many types of bacteria, sucrose is hydrolyzed to glucose and tose and then utilized by these organisms to form many different commods, many of which are capable of reduction and some of which are ie. The reducing conditions resulting from the organic compounds ned protect hemoglobin and myoglobin from irreversible oxidation of iron to the ferric state and insure the formation of nitric oxide nitric oxide hemoglobin. During storage these reducing conditions perand the chances that the nitric oxide hemoglobin will form methemonin are minimized. However, when exposed to oxygen, nitric oxide oglobin is oxidized to methemoglobin. Thus sliced bacon turns brown tanding for some time.

nce sucrose forms glucose and fructose, the effects of other monodisaccharides have been tested. For the most part they are not as sucred in the pickle as sucrose because they are used by the bacteria apidly that the pH falls to a range where methemoglobin formation reeded up and instead of protecting against browning these sugars can hasten it. In short cures they are useful, and today glucose in the 1 of corn sugar or a mixture of glucose is sometimes used along with ose.

king Meats

ardwood smokes are known to produce superior preservation and or. Jensen²⁶ lists the effects of smoking: (1) drying effect. (2) impart-lesirable organoleptic properties, (3) bringing out color inside the cured :, (4) imparting antioxidants to the fat, (5) impregnating the outside tons of the meat with constituents of smoke that serve as antiseptics germicides, (6) an adjuvant action of constituents of smoke and heat nicroorganisms, if the process is carried out at a temperature above (120°F), (7) a tendering action from the high humidity of the smoke-em combination with its high temperature, (8) a diminution of nitrite and probably by reason of the aliphatic diazo reaction with protein hoccurs at higher temperatures, and (9) the imparting of a desirable or gloss on both the skin and the flesh sides of meat through the interpretation of the piece.

take contains many compounds produced by fracture of the large cules present in the wood. In destructive distillation of wood, the large taken the large and these compounds are distilled out and the large taken to the large taken to the large cules present in the wood. In destructive distillation of wood, the large taken to the large cules present in the wood, and these compounds are distilled out and the large cules are t

TABLE 6.2. HARDWOOD SMOKE

	Parts Per Million
Formaldehyde, HCHO	25-40
Higher aldehydes	140-180
Formic acid, HCOOH	90–125
Acetic and higher acids	460-500
Phenols	20-30
Ketones	. 190–200
Resins	over 1000

separated. However, the conditions during smoking are more conbecause not only are these compounds formed and distilled out of but, in the presence of fire and air, they may burn or react vanother. In the destructive distillation of wood sizeable quar phenols are formed, for example, but in smoke the concentrations because they react with formaldehyde and form resins. Jensen a following composition of hardwood smoke based largely on work and Lane. 40 (See Table 6.2.)

The research laboratories of many meat packing houses have stubactericidal effect of smoking and have found reduction in numbacteria. The concentration of formaldehyde in smoke is sufficien to account for this effect and some investigators believe that active ingredient. There is the possibility that other compound smoke such as some of the higher aldehydes may augment this effect temperature of the smokehouse is usually sufficiently high so that the bacteria are destroyed by heat. Smoked meat has a marked included heat in the presence of the meat and act as bactericides. It is since molds are much more tolerant of aldehydes than bacteria, has little effect in preventing the growth of a number of type many molds will grow in the presence of sizeable quantities of though molding is sometimes a problem with smoked meats, none molds produce disease in man.

The gloss formed on the surface of smoked meats is the result effects. Resins formed in smokes from the reactions of aldehydricularly formaldehyde and phenols—are deposited on the surface meat. Also at the temperature of the smokehouse some fat oozes a sen reports that the aldehydes produce a glossy finish when the roiled out a little.

During smoking there is a steady decrease in the amount o Nitrous acid reacts with free amino groups to give the aliphat reaction and form hydroxyl groups.

$$-NH_2 + HONO \rightarrow -OH + N_2 + HOO$$

when meat is smoked, the onset of rancidity is delayed. The protection marily on the outside of the smoked piece since there is little penetraof the smoke compounds to the center of a ham or bacon. When the
cas sliced, it is for the most part unprotected. The identity of the
pounds that act as antioxidants has not been established; but since
if amounts of phenols are in the smoke and since many antioxidants
whenols, it has been assumed that these are the active agents. Watts
orts her work with Faulkner on the effect of liquid smoke as an antilint. Liquid smoke is produced by the thermal decomposition of hardds and a number of these products are on the market. They are not
wed by the Bureau of Animal Industry in commercial production of
ked meat. Watts and Faulkner found considerable difference in the
oxidant power of these liquid smokes, ranging from no effect at all to a
nounced inhibition of oxidation.

The reaction is not completely known, but it is believed that nitric is myoglobin is denatured and forms the corresponding porphorin and stured protein. The solubility is diminished on coagulation and the insecd stability or decreased reactivity may be the result of this change.

the moist heat of the smokehouse the muscle proteins are greatly cted. Denaturation of the protein occurs at the higher temperatures. At beginning of the smoking process when the hams are warmed, autolytic in es in the cells may be active and cause some tendering. As the inal temperature of the piece of meat increases, enzymes will also be dered and cease to function. This will be particularly true of meat ked to a high temperature or held in the smokehouse for a relatively time. During smoking the changes that occur with cooking get under and when the time of smoking is long or the temperature high, as a timple in the preparation of ready-to-eat ham, these changes go to pletion.

ANGES IN MEAT ON COOKING

ooking causes many changes to occur in meat which develop appetiteul ting flavors. Who has not smacked his lips over the thought of a no n steak dripping with juicy goodness? Yet the sight and odor of ame steak before it is cooked inspires none of the same response in doubtedly, cooking markedly increases those properties which are ad organoleptic.

to apply as far as our well being is concerned, the most important

change in cooking is the destruction of microorganisms. All cooking esses decrease the number of bacteria, yeasts, and molds, and if the m well done may even render it sterile. Spore-forming organisms ar stroyed and the spores killed if the meat is cooked to a sufficiently temperature. In canning meat this is important since the meat must be pletely sterile so that it will keep. Pork is often infected with a par *Trichinella spiralis*, which is dangerous to man. This organism is at 132°F; Jensen points out that the Bureau of Animal Industry den that pork be cooked to an internal temperature of 137°F to insure, the 5° margin of safety, that all organisms are destroyed.

The chemical and physical changes which occur on cooking ar merous and although some studies have been made on this problem the actions are not all known or understood. They are: (1) denaturated the protein, (2) hydrolysis of collagen to gelatin, (3) color change formation of drip, (5) development of brown flavor, (6) rupture of fat and dispersion of fat through the meat, and (7) decrease in vitamins possibly decrease in the nutritive value of the protein.

Denaturation of the Protein

This is evidenced by the change in the physical condition of the r The jellylike structure stiffens and often toughens. There is a shrin in volume and an increase in density as cooking proceeds. Ramsbo and Strandine⁴¹ studied the change in tenderness of various muscles wholesale cuts of beef of three young heifers, good grade, by the test. This is a test in which tenderness is determined by measuring force necessary to cut through a standard sample. They found that tel ness decreased in 35 of 50 samples cooked in lard to an internal temp ture of 170°F. Lowe³² discovered in the case of beef rib roasts that meat is tenderest, rare is intermediate, and well done, least tender. used both the shear test and a penetrometer. Other workers have for similar results, but some have found divergent ones. Undoubtedly the ference in results is caused by the fact that a measure of tenderne meat does not measure the change in tenderness of the protein. Ins a change in the tenderness of meat is the result of numerous cha only one of which is this change in protein. Heat denaturation of pr is a familiar experience in food cookery, but it is not always accompa by a toughening of the protein.

Denaturation of a protein molecule is believed to be a series of tions in which the protein molecule is altered by splitting of some o links, particularly some of the hydrogen bonds and sulfur-sulfur which help maintain the three-dimensional shape of the molecule. The alar weight of the molecule may not be very much changed in the first as of the reactions. In heat denaturation, as in other types of deuration, it is believed that these changes in the molecule account for alteration in such properties as solubility and density.

Hydrolysis of Collagen

his is the reaction by which commercial gelatin is prepared. The exto which it occurs in meat cookery has been the object of a number esearches. If a piece of meat is cooked for a long time, the loose nective tissue disappears and the gravy gels on cooling. This is parlarly noticeable in moist cooking such as braising. However, even in cooking such as roasting and frying, the connective tissue and the agen fibers are in contact with water from the tissue. Connective tissue If has a fair amount of water. So there is always the possibility that rolysis of the large collagen molecules to the relatively small gelatin lecules can occur.

Vinegarden, et al. carried out careful tests of the connective tissue m nine beeves. They used the ligamentum nuchae in which most of the ers are elastin, the deep flexor tendon in which most of the fibers are agen, and strips from the aponeurotic sheet which is intermediate in agen and elastin content between the first two tissues. Collagen and al nitrogen were determined on all the samples. The strips were heated listilled water for periods of time of 1, 2, 4, 16, and 64 minutes and emperatures ranging from 60°C to 95°C. Changes in shear force, ght, length, width, and thickness were measured. The tissues were also died histologically. Heating for 64 minutes at 95°C caused some changes the collagenous fibers, so that some were fused or merged, some rightened, and all were less distinct. The elastin fibers did not show se changes. All tissues showed softening and a decrease in shear force, n the ligamentum nuchae, on heating at the higher temperatures. There . little change at 60°C but the extent of change increased with inused temperature and increased time; 60°C or 65°C is the internal temature to which meat is cooked when it is rare. Thus the study indicates t although little change in collagen can be expected for rare meat, agen hydrolysis increases with doneness. The study also demonstrated t strips from meat aged 35 days had slightly lower shear forces than e ared 10 days; finally the older animals for the most part required e force for shearing the tissue even though tissues from the oldest ani-1. in 8-year-old cow, did not have the highest shear-force readings.

reswold! studied a number of factors in beef rounds, commercial and ne grades, cooked by various methods. Collagen was significantly

higher in the raw meat for the commercial grade over the prime but not the top round over bottom round. The loss of collagen on cooking curred with all methods and increased with an increase in the internal perature to which the round was cooked. When pressure cooking was opared, it was found that meat cooked at 10 lbs or 15 lbs pressure sho greater losses of collagen than that cooked at 5 lbs pressure. It was high in meat roasted at 250°F than in that roasted at 300°F or braised. No nating in vinegar caused a greater loss; but other methods such as scopounding, or using enzyme preparations had little effect.

Color Change

The color change from red or purplish red to brown or gray on coo has already been discussed under the color of meat. When meat is coo oxyhemoglobin and oxymyoglobin (red) and hemoglobin and myoglo (purplish red) are denatured. The ferrous iron in the free porphyrin for is rapidly oxidized to ferric iron of hemin (brown). Some transformatof oxyhemoglobin, oxymyoglobin, hemoglobin, and myoglobin to rhemoglobin and metmyoglobin (brown) can also occur.

Drip Formation

This accounts for some of the change in weight that occurs in a piece meat during cooking. The drip is composed of water carrying a number soluble compounds as well as some coagulable protein and fat. As cook proceeds the coagulable protein is denatured by the heat and forms a continuous in the pan gravy. Numerous compounds have been isolated from the debut it is questionable whether they account for the delicious flavor of mixture. Most of the compounds are small molecules formed either durautolysis while the meat is ripening or by heat fragmentation of lating molecules. Creatine, creatinine, salts, particularly sodium chloride, stramounts of amino acids, amino acid derivatives such as amines, puring and pyrimidines can be detected. These molecules are individually flavoless or almost so aside from the saltiness of sodium chloride and of salts.

Fat oozing always occurs but the extent to which fat runs out of the medepends on many factors. The cut of meat and the amount of fattiness, method of cooking, the extent to which the piece is cooked all obvious affect the amount of fat in the drip. There may also be fat degradate products in the drip. When steak is broiled and some burning of the fat curs, small amounts of acrolein can be detected in the drippings. Free facility are sometimes present. Griswold¹⁹ found that the amount of solunitrogen compounds increased in the drippings during cooking but did

that free amino nitrogen showed an increase. She cooked round steak various methods.

at Aroma

he change in flavor that occurs with cooking is one of the most apmit changes to the man at the table. Particularly where the meat has uned, a strong flavor, sometimes called the "brown flavor," is evident, cker in a review points out that raw meat, whether beef, pork, lamb, hicken has little flavor. It is slightly salty and slightly sweet. Cooked t likewise has this slightly salty, slightly sweet taste but it also poses an aroma that adds greatly to the flavor. The aroma is composed of molecular weight, volatile compounds such as amines, ammonia, hygen sulfide, and organic acids. These compounds probably arise from cracking of amino acids during heating. The reactions may be decarylation, deamination, or desulfuring in which either the free amino acid olypeptides react. The compounds formed vary according to the species, hat the flavor of lamb is recognizable and different from the flavor of f.

persion of Fat

at cells rupture and fat disperses through the meat on cooking. The teins in fat cells of the adipose tissues undergo denaturation on cookjust as all other proteins in the meat. There is a change in the permetity of the cell walls and fat flows out. Wang, et al. studied this nge carefully in beef steaks. They used the longissimus dorsi (ribeye) and semi tendinosus, eye of the round, and broiled them to an internal tempature of 150° F. Sections made of the cooked steak were stained with an IV or Nile Blue to show the fat. It could be seen that fat had diffed out of the fat cells of the perimysium. Fat droplets seemed to be dissed along the path of degraded collagen and mixed with it. The size of droplets decreased from the dispersion center to the periphery. Wang, I interpreted this to mean that the degraded collagen emulsifies the fat. stographs of their slides are shown in Figure 6.5.

crease in Vitamins

The decrease in vitamins that occurs during cooking of meat has been exvery studied. The B complex vitamins are all more or less sensitive to toll and thiamin and pantothenic acid are particularly labile. If cooking relonged and if the temperature is high, the destruction of the vitamins 3 be appreciable. There is always some loss. Niacin and riboflavin are



1. Raw fresh light go control, showing par perimysial fat islated cells intact and loaded × 20.

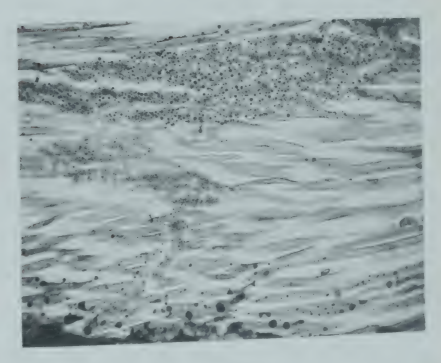


2. Section of above sample it was cooked, showing depleted fat islands sure by partly hydrolyzed of Three empty fat cells with tinct walls and nuclei a in upper fat island, more than half of the the lower island have their fat. × 55.

FIGURE 6.5. CHANGES IN MEAT FAT ON COOKING. Fat is stained and appears dark.

ther cooked ribeye, showtypical fat dispersion center, spersed fat has followed a arse approximately coincidwith that of hydrolyzed lagen with little or no fat in domysial spaces. × 55.



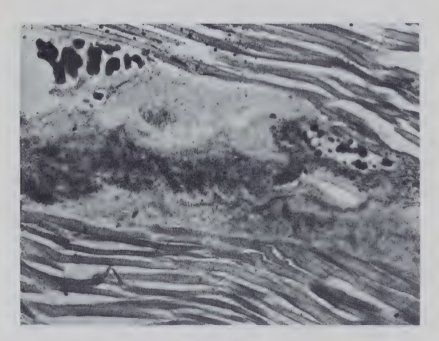


other cooked ple showing ensive dispern from fat a not shown, above upper

FIGURE 6.5 (Continued)



 Cooked 14-day-aged p eye, with dispersion of neighboring perimysia which separate muscle × 55.



6. Cooked fresh prime ribeye perimysium contains thoroughly mixed fat and hydrolyzed collagen. Note relatively intact collagen fibers between 2 fat islands are free of fat. × 55.

FIGURE 6.5 (Continued)

coked unaged cow round. With intact (right) and hyrolyzed (left) collagen. Fat sland at lower right. × 110. oduced from Wang, H., h. E., Bates, U., Beard, F. J., c., J. C. and Hankins, O. C., Research, 19, 314-22 (1954). tesy of American Meat Inferoundation.



RE 6.5 (Concluded)

the to heat and do not disappear as rapidly during the cooking of the theorem, the folic acid group is very sensitive to heat, and as much oper cent may disappear. Ascorbic acid begins to decrease as soon as nimal is slaughtered, and during cooking any that is still present will be further diminution. Cooked meat cannot be counted on to supply the than very small amounts of ascorbic acid to our diet.

face Reddening

consionally a red surface is produced when meat is boiled, broiled, or ted and sometimes it is seen in canned meat. We do not expect meat to red when cooked, but instead anticipate a brown or gray color and, ared meats, a pink. Jensen discusses the conditions under which it is the for the reddening to occur, but points out that sometimes it is produced under conditions where as yet there is no explanation. The reddening and make the meat harmful or unpalatable. A red color is produced of the meat comes in contact with nitrites, nitrates, carbon monoxins. If the meat comes in contact with nitrites, nitrates, carbon monoxins. If the mounts is altitle as a few parts per million of nitrite can cause the reddening as nitric oxide hemoglobin or the porphyrin is formed.

trates. If meat is cooked with these vegetables, some leaching of to can occur and reddening of the meat surface will form. Carbon mode might come in contact with roasting meat through a leak in a gast carrying coal gas (natural gas has little or no carbon monoxide), carbon monoxide is a very poisonous gas, neither carbon monoxide globin nor carbon monoxide myoglobin are harmful. Sulfites can also duce reddening, but they are not likely to be present in either the not vegetables. Sulfites are prohibited in meat and do not occur in platterial.

Overcooking

This causes an excessive loss of drip and a toughening of the mean mean becomes stringy as the amount of connective tissue falls and the drips out. Not only are the tenderness and juiciness of the mean diminishment the flavor as well decreases. Volatile, odorous compounds that tribute to the flavor are driven off during the excessive cooking period the mean becomes less and less flavorful. A decrease in the nutritive value protein may also occur (p. 136). Overcooking can be the result eit too long a time or too high a temperature.

TENDERNESS

Tenderness is one of the most important attributes for rating me we bite into a piece of meat, tenderness is one of the first sensation ceived. Like many other characteristics of meat, tenderness is incompunderstood. However, we are fairly certain that it is the result of a plex of factors. Most investigators find that tenderness of meat decon cooking. Indeed, some restaurants print on the menu that tenderness teaks cannot be guaranteed unless they are served rare. But since coincludes a whole series of changes, these observations do not expla fundamental cause of tenderness or its absence.

Tenderness is measured by one of three methods or a combinate them: (1) the force necessary to penetrate through a piece of meat, (shear test which measures the force necessary to cut across fibers, or panel of trained judges. The correlation between methods is sometime cellent and sometimes poor.

The factors that may be significant in explaining tenderness are (amounts of connective tissue in the muscle, (2) the amount of hydrat the muscle proteins, (3) the nature of the protein molecules, and (amount of fat between muscle fibers. These factors will be discusse evaluated.

nnective Tissue

The connective tissue that forms a fine network through muscle tissue been credited traditionally with tenderness or lack of tenderness. Howfinding methods by which connective tissue can be estimated without and separated or differentiated from other types of tissue has been cedingly difficult. Investigators have for the most part used estimates of ugen, the protein present as fibers in the ground substance, to measure nective tissue. It has been claimed that collagen is completely hydrod to gelatin by autoclaving and that it is the only protein in a meat ple capable of this reaction. A number of determinations of the ount of nitrogen in the gelatin formed have been presented as an estie of the connective tissue present. Some years ago it was realized that assumption that only collagen would be hydrolyzed is problematic. the method was then refined by preparing the meat sample by extracbefore autoclaving. In this modification it is assumed that the proteins able of some hydrolysis on autoclaving are first removed and that none he collagen is removed. Miller and Kastelic have critically studied ne of the methods used to evaluate connective tissue in bovine muscle fnd that the amount of protein extracted by alkali increases with time sposure of the sample and with the concentration of the alkali. (Sodium rexide is usually used.) The protein extracted probably is from the and substance and not collagen since the protein is free of hydroxyline. Their work shows that nitrogen determinations on insoluble subces do not give valid measures of collagen or elastin.

he hydroxyproline method for estimating either collagen or elastin deds on the presence of unusually large amounts of hydroxyproline in a proteins. However, the proteins used in setting up the method are present in tendons, hide, and ligaments. When the method is applied neat, collagen is calculated from the hydroxyproline determination on gelatin formed by autoclaving, while elastin is calculated from a demination of hydroxyproline in the portion insoluble in alkali and after pelaving. Elastin is assumed to be resistant to hydrolysis by mild alkaling water on autoclaving. Sometimes the residue remaining has been sured by estimating the nitrogen in it and calculating this as the elasting ion.

Samples of meat are prepared on slides and stained so that the contre tissue can be seen and evaluated. This is very difficult to do prey. Harrison, et al. 22 demonstrated that the most tender piece of beef d 1, 2, 5, 10, 20, and 30 days) had the smallest amount of connective Deatherage's group²⁴ estimated the amount of collagen and elastin by termining the alkali insoluble fraction. Their beef, short loin, aged 3 c showed no correlation between this fraction and tenderness but when beef was aged 15 days, there was a definite correlation. In another series beeves representing a wide variety of market grades, they²⁵ found exce correlation between alkali insoluble protein and tenderness. These ani had all been aged 14 days at 3.5°C.

So far no simple and reliable method has been developed for estimate connective tissue. The connective tissue in muscle has not been separate difficult job, and the amount of collagen and elastin determined. The lation of connective tissue to tenderness is consequently still not content with the work of the amount of connective tissue can not be related with tenderness in all cases.

Hydration

The degree of hydration of muscle proteins may influence tenderness migration of ions occurs² as the meat ages. Calcium, sodium, and mag sium migrate out of the muscle cell while potassium ions move in. The texchange is such that the over-all change is an increase of ions in the petein. These ions are held so firmly that they not only do not appear in juice formed when the meat is cooked, but they also are not extracted water. Deatherage concludes that the migration of ions allows greater dration of the protein and consequently an increase in tenderness. This logical conclusion.

Water of hydration is held within a protein molecule by hydrogen bo ing as the slightly positive hydrogen is attracted to relatively nega atoms of oxygen or nitrogen. Water molecules are held between the fo or coils as well as on the surface of the large protein molecules. Hyd tion is affected by pH and is lowest at the isoelectric point of the prot Ions also influence the ability of the protein to hydrate—hydration to ing to increase with increase in ionization. During rigor mortis, the pH carcass of a well nourished animal falls to approximately 5.6, very clos the isoelectric point of actomyosin; and the amount of hydration decrea This is demonstrated by the increase in the amount of juice formed w meat in rigor is cooked. As rigor passes the pH rises slightly, so that protein begins to form negative ions rather than the internal zwitter: of the isoelectric point. The migration of ions allows salt formation the protein may become more highly ionized and more able to hold w molecules by hydrogen bonding. Deatherage believes it is reasonable to pect a highly hydrated protein to be more tender than one which is no by hydrated. The protein molecules will separate from one another re-readily if a layer of water lies between them.

ture of Protein Molecules

there are some observations which suggest that changes in muscle prosinfluence tenderness. Harrison, et al. studied the histological appearance of sections of the psoas (tenderloin) muscle in beeves of various derness. In the most tender specimen there was an absence of longinal striations on the muscle fibers; whereas, in the least tender a cimen from a cow 8 years old and of cutter grade—the fibers were sted and gnarled, less uniform in diameter, and the cross striations less form in spacing. This observation indicates that protoplasm protein, contractile material of the cell, is involved.

Two proteins, myoglobin and actomyosin, have been studied by Deathge's group. Myoglobin content gave a positive correlation with tenders in rib steaks which had been aged 15 days. They found no correlator actomyosin content with tenderness. As rigor occurs and the muscle omes rigid and tough, actin and myosin combine to form actomyosin, rigor passes and the meat begins to become tender, the content of actin myosin continues to fall. The change in the muscle is not, therefore, a spie reversal of actomyosin formation. Likewise water extractible nitrowhich includes polypeptides and peptones, the fragments of proteins, as not increase in amount but instead falls.

Deatherage and his group have studied a number of other cell comvents and have shown that there is no correlation of tenderness with al nitrogen, trichloracetic acid-soluble nitrogen, nonprotein nitrogen, er-soluble nitrogen, pH, lactic acid, moisture or inorganic phosphate.

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A number of investigators believe that good marbling contributes to derness of meat. Harrison et al.²² noticed one or two rows of fat cells ween the fibers in their most tender specimens and Husaini found a coration, although of a low order, between tenderness and intermuscular The effect of marbling on tenderness is still open to question.

ICINESS

The amount and the quality of the juice formed when meat is chewed is ther of the most important factors in judging the total quality of a ce of cooked meat. But it is a factor very difficult to define. Everyone experienced the extremes of a juicy bite of meat or a very dry one, but problem of exactly defining juiciness or setting up some device that can

objectively measure it has been extremely difficult. Several instrument measuring press fluid—the amount of juice that can be pressed o meat sample under high pressure—have been devised and used, correlation of the results of these instruments with the judgemen panel have not been good. In general it has been considered that m of the meat with fat, is one of the most important factors in produc sensation of juiciness. In one extensive study, Gaddis, Hankin Hiner¹⁶ found a low but significant correlation between the amount of fluid and the juiciness scores in 97 beef ribs, but none in 115 lam sheep (legs studied) or 11 goats. With beef, increase in percentage o the press juice caused a decrease in the amount of juice yet an incr the juiciness scores by the panel up to 2 per cent fat. They conclude the sensation of juiciness is probably the result of numerous factor one of which is the juice in the piece of meat. They believe that fa flavor and stimulates the flow of saliva in the mouth. It also coa mouth and prolongs the sensation of moistness and richness duri chewing and swallowing. These combined effects give the sensation o ness.

PHOSPHATES IN CANNED MEAT

The effect of phosphates on the ability of meat to bind water has investigated in recent years. Some results appear to indicate that the is specific for phosphate, others that it depends on the increased pH occurs. Hamm and Grau²¹ found that the addition of one per cent phate to the sodium chloride that is put in meat has a definite eff the amount of bound water and decreases the loss of drip. Bendall⁵ s the effect of orthophosphate, pyrophosphate, metaphosphate, Calgorian commercial polyphosphate), and phosphate glass on rabbit meat. Alt all influenced meat's capacity to bind water, the pyrophosphate has greatest effect. Bendall considers that the binding of water depends ionic strength present except in the case of pyrophosphate.

Commercially, phosphate is now added to canned meat as well shrimp. The increased binding of water decreases the shrinkage oprocessing and improves tenderness.

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Vegetables and Fruits

Webster's International Unabridged Dictionary says, "There is well-drawn distinction between vegetables and fruits in the popular set but it has been held by the courts that all those which, like potatoes, of bage, carrots, peas, celery, lettuce, tomatoes, etc. are eaten (whether coor or raw) with the principal part of the meal are to be regarded as vegetable while those used only for desserts are fruits." Botanically, fruits are considered the ripened seeds and the adjacent tissues which contain them at this definition is often carried over into foods. So apples, peaches, banar cherries, and berries are always called fruits while tomatoes and melons sometimes classed one way and sometimes another. Cucumbers, pepp squash, and eggplant are usually classed as vegetables, although they botanically the "fruit of the plant."

Fruits and vegetables add enormously to the interest and variety of diet by their great range of color and texture and by their complex aron giving each variety and even each individual a slightly different flavor. It tritionally, fruits and vegetables are important because they contain la amounts of certain vitamins and minerals. Cooking often brings about change in the aroma and even in the relative sourness and sweetness, through the use of fresh and cooked fruits and vegetables, the possible viations in menu pattern are immense.

Although fruits and vegetables vary greatly in their chemical competion, some generalizations are possible. All, with the exception of nuts dates, are high in water with a range from approximately 70 per cent pears, bananas, figs, etc. to 98 per cent for vegetable marrow. All, with exception of legumes and nuts, are relatively low in protein. Although quantity of protein in vegetables and fruits is sometimes neglected, no cular material ever exists without a certain amount. Protein varies from proximately 0.3 per cent in apples to 4.4 per cent in brussels sprouts.

Il vegetables and fruits contain some carbohydrate. Part of the carborate in fresh fruits is present as cellulose and pectic substances in the walls, but these compounds are indigestible and not available to the tan body. Starch is present in almost all fruit and vegetables although it disappear on ripening. Glucose, fructose, and sucrose are widely disted, and sweet taste is dependent on their occurrence. Glucose, fructose, and starches constitute the "available carbohydrate" of and vegetables, and the caloric value of the food depends in large sure on the concentration of these components. In practical dietetics and vegetables are frequently classified by the approximate amount vailable carbohydrate since all those in the same class have approxicly the same number of calories. The common classifications are: 5, 10, and 20 per cent available carbohydrate. The amount of lipids in fruits vegetables is usually very small. Nuts are the general exception, while we vegetables such as avocados are a rich source of fat.

GETABLES

Leaf vegetables (lettuce, mustard greens, chard, spinach, water cress, sley, cabbage) are high in water and cellulose and low in calories and tern. They add valuable amounts of minerals and vitamins to the diet alugh they do not contain large amounts of most of these nutrients. They usually rich in iron and provitamin A, and often in the B complex. The leaves often contain appreciable amounts of ascorbic acid.

Flowers, buds, and stems (broccoli, cauliflower, artichoke, asparagus, ry, kohlrabi) are relatively high in water and cellulose, but low in pro-

They have moderate amounts of calcium and some are moderately in provitamin A. They all contribute small amounts of vitamins and erals. A few have moderate amounts of ascorbic acid and riboflavin.

Bulbs, roots, and tubers (potatoes, beets, turnips, carrots, rutabaga) high in water, moderate in cellulose, and contain an appreciable ount of available carbohydrate. The available carbohydrates are ches, glucose, and some sucrose. The amounts of the vitamins and mins are not high but are valuable adjuncts to the diet. The amount of orbic acid in potatoes has often made the difference between gross vv and its absence in large numbers of Europeans.

seeds (legumes, corn, rice) are relatively low in water and cellulose, amount of protein and a large amount of starch. They are

tole sources of the B complex vitamins and iron.

Wegetable fruits (cucumbers, peppers, melons, tomatoes, squash, opkin, eggplant, okra) are relatively high in water and cellulose but low alones and protein. Many contain valuable amounts of vitamins and

small amounts of minerals. Some, such as the tomato, are notable for corbic acid; others, such as green peppers and squash, for provitami and some, such as the tomato, for thiamine.

STRUCTURE OF FRUITS AND VEGETABLES

A fresh fruit or vegetable is a group of living cells, still undergoing m bolic reactions. Although when harvested it is cut off from tissues of plant supplying water and other nutrients, it is nevertheless still living. chief type of cell in the edible portion of most fruits and vegetables is parenchyma cell. See Figure 7.1. It makes up the bulk of the cells in lea fruits, and even in young edible stems. A parenchyma cell is rather walled; it may be polygonal or cubical in shape, but all are about the s size. The parenchyma cells do not fit tightly together but are often arated by air spaces that contribute to the slightly chalky appearance fresh fruit or vegetable. The walls of parenchyma cells in young plants composed almost entirely of fibrils of cellulose. The cells are held toge by cementing substances, which in the young plant are composed of pe substances. As the plant grows older the nature of these cementing s stances often changes, lignins and other compounds are deposited, and cellulose layer of the cell wall thickens. Such an old plant is not d sidered desirable for food since it is woody and tough even when cooked

The material within the cell wall is protoplasm composed of a very lanumber of different molecules that form either a viscous fluid or a Some of the molecules are dissolved in water in the protoplasm, but m such as proteins are colloidally dispersed. The protoplasm is not unifor but differentiated into various regions and cell parts, the most distinct which is the nucleus. It is believed that much of the activity of a cell directed by the nucleus and that a cell cannot survive long without Within the cytoplasm are numerous small bodies called plastids. Parenchyma cells of leaves and occasionally other tissues contain grant green plastids called chloroplasts—the site of the chlorophyll so import in photosynthesis by the plant. If the plastid is any other color because the occurrence of other pigments in it, it is called chromoplast. The let plasts are colorless bodies containing starch granules.

Beside the plastids there are often large vacuoles in cells, made up droplets of solutions with strands of cytoplasm around them. They consalts, sugars, and other soluble material dissolved in water. This solution sometimes spoken of as the "cell sap." In young cells the vacuoles small and numerous. As the cell grows the total size of the vacuoles creases much more rapidly than does the amount of protoplasm, through the imbibition of water and other small molecules. The vacuoles coaled

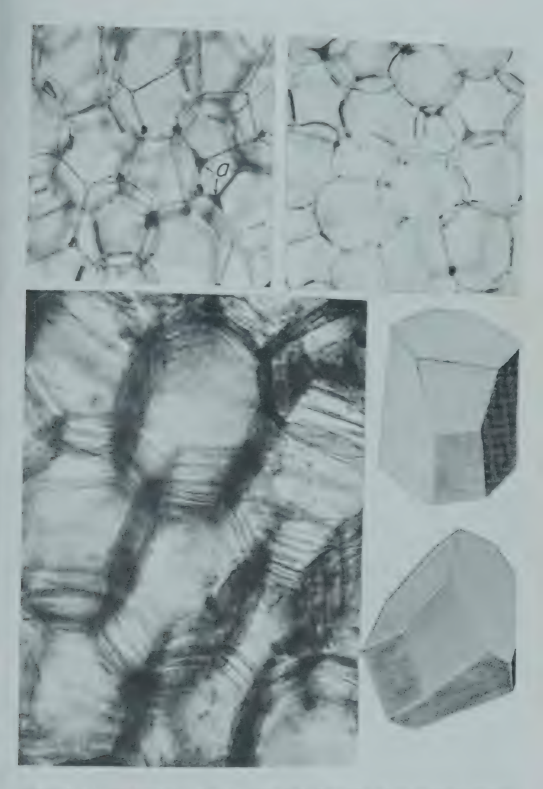


FIGURE 7.1. PARENCHYMA CELLS. Upper left: From living onion: dark regions, (a) are intercellular spaces filled with air. Currier, 1946. Upper right. From blanched onion, intercellular spaces are filled with water. Currier, 1946. Lower left. Nonliving, with much intercellular air. Lower right. Two diagrams, showing three-dimensional shape. Marvin, 1939.

Courtesy of Elliot T. Weier

and become much larger in size and smaller in number with often only large vacuole in a mature cell. Cells with a high fat content may also an oil vacuole. Some cells contain crystals embedded in the cytoplasm.

Other types of plant cells besides parenchyma cells are the conduction cells, the supporting cells, and the protective cells. See Figure 7.2. conducting cells are composed of long tubes through which water and or foodstuffs are distributed through the plant. These are of two to called xylem and phloem. The walls of the xylem are composed primarically cellulose thickened at intervals in definite patterns with lignin. The work of the phloem contain little lignin. Fibers composed of cellulose may of associated with the phloem. When numerous or large, such fibers are jectionable since they are largely unchanged on cooking and procestringiness and toughness.

The supporting tissues are not numerous in young plants or in the yo parts of plants, desirable for foods. They are composed of long poin cells whose cell walls of cellulose thicken as the plant ages and becomercusted with lignin. Some plants have another type of supporting ce which the cell wall is composed of cellulose and pectic substances in p of cellulose and lignin.

The protective tissue is composed of specialized parenchyma cells that crete cutin or contain suberin. Sometimes these cells are thick and corky other plants they are thin. But they are closely pressed together and usually quite tough. Usually the epidermis of a fruit or leaf contastomata, minute valves through which exchange of gases can occur we they are open. The cutin or suberin of the epidermal cells make them pervious to water and also protect them from injury. Often this layer cells forms a skin or peel which may be removed in the preparation of leaf or fruit for eating. The layer of epidermal tissue not only protect the organ from mechanical injury, but also prevents inroads by insefungi, and microorganisms. Everyone has observed that after the skin broken the keeping time of a fruit is relatively short, whether it has beharvested or not.

The cells of a harvested fruit or vegetable are still *living*. It is only w food is cooked that cells are killed. If the food is frozen or kept for a l time, death of cells usually occurs. See Figure 7.4.

The composition of the cutin or suberin of plants is for the most punknown. A number of studies have been made of the composition waxes such as carnauba or japan wax which are produced commercia and some investigation has also been directed to discovering the composition of the cutin of other plants, but not necessarily those which used for food. The cutin of a few food plants, particularly apples, has been directed to discovering the composition of the cutin of a few food plants, particularly apples, has been directed to discovering the composition of the cutin of a few food plants, particularly apples, has been directed to discovering the composition of the cutin of a few food plants, particularly apples, has been directed to discovering the composition of the cutin of a few food plants, particularly apples, has been directed to discovering the composition of the cutin of a few food plants, particularly apples, has been directed to discovering the composition of the cutin of a few food plants, particularly apples, has been directed to discovering the composition of the cutin of a few food plants, particularly apples, has been directed to discovering the composition of the cutin of a few food plants, particularly apples, has been directed to discovering the composition of the cutin of a few food plants, particularly apples, has been directed to discovering the cutin of the cutin of a few food plants, particularly apples are discovering the cutin of the cutin of



or the leaf blade. Lower left: Cross section of an old (inedible) asparagus—conducting and strengthening tissues a are grouped in discrete bundles and surface by parenchyma cells b, fibers have formed just under the epidermis of the stem c. young asparagus stem there would be no fibers and the bundles would be in a rudimentage of development. In many stems the conducting cells form a continuous cylinder n the stem.) Lower right: Longitudinal section of old asparagus stem showing the conng and strengthening bundles a embedded in parenchyma tissue b.

Courtesy of Elliot T. Weier.

ionated and the components at least partly identified. The composition ost waxy coatings is complex but the compounds fall into a few general es. The components of natural waxes have been classified in the folgategories: (1) hydrocarbons, (2) fatty and wax acids, (3) ketones, and wax alcohols, (5) esters, (6) ethers, (7) pseudo esters, and (8) ratic compounds. These compounds usually have long chains of benut and 22 carbons or more for the principal chain. The principal acids, palmitic, stearic, oleic, linoleic, and linolenic, are found in

many of the waxes as esters, as well as free acids. Some waxes contain acids with a higher number of carbons in the chain than the more confatty acids. Thus it is reported that carnauba wax contains 75 per myricyl cerotate.

Cerotic acid is the straight chain fatty acid with 26 carbons. A sum is presented in Table 7.1.

TABLE 7.1. COMPOUNDS REPORTED IN PLANT WAXES

No. of Carbons		
	Alcohols and Ke	tones
12	1-12 dodecandiol	CH ₂ OH(CH ₂) ₁₀ CH ₂ OH
25	Pentacosan-8-ol (or -one)	$CH_3(CH_2)_{16}CHOH(CH_2)_6C$
26	Hexacosanol	$C_{26}H_{53}OH$
27	Heptacosan-9-ol (or -one)	$CH_3(CH_2)_{17}CHOH(CH_2)_7C$
28	Octacosanol	C ₂₈ H ₅₇ OH
29	Nonacosan-10-ol (or -one)	$CH_3(CH_2)_{18}CHOH(CH_2)_8C$
29	Nonacosan-15-one	$CH_{3}(CH_{2})_{13}CO(CH_{2})_{13}CH_{3}$
30	Triacontanol	$C_{30}H_{61}OH$
31	Hentriacontan-16-ol (or -one)	$CH_3(CH_2)_{14}CHOH(CH_2)_{14}C$
33	Tritriacontan-16-ol (or -one)	$CH_{3}(CH_{2})_{16}CHOH(CH_{2})_{14}C$
	Acids	
8	Suberic Acid	COOH(CH ₂) ₆ COOH
12	Sabinic Acid	$CH_2OH(CH_2)_{10}COOH$
20	Eicosandicarboxylic Acid	$C_{18}H_{36}(COOH)_2$
22	Phellonic Acid	CH ₂ OH(CH ₂) ₂₀ COOH
26	Cerotic Acid	C ₂₅ H ₅₁ COOH
26	Juniperic Acid	$CH_3(CH_2)_9CHOH(CH_2)_{14}CO$
30	Ursolic Acid	CH ₃ CH ₃
	H	CH ₃ CH ₃ CH ₃ CH ₃ CH ₃
	Hydrocarbon	ns .
27	Heptacosane	$C_{27}H_{56}$
29	Nonacosane	$C_{29}H_{60}$

ott" has reported histological studies on the cells of numerous plants one that are used for foods. She found that the walls of all intercelluir spaces in leaves, stems, fruit, and flowers are lined with a substance t does not dissolve in 80 per cent sulfuric acid or in chromic acid when ssue section is flooded with the reagent. She calls the substance or mix-"suberin" and indicates that it is waxy. She also found the material in n layer on the inside of cells next to the protoplasmic membrane in hairs and along vessel segments. The degree to which this material ocvaries with the type of plant and increases on ageing. The material be stained. Usually the term "suberin" is only applied to corky tissues. ius we see that many plant tissues are covered with a water-impervious of waxy material called "cutin." The cutin limits the amount of water by transpiration, as well as the amount of water-soluble substances lost ne leaching of rain. The correlation between the thickness of the whole le layer and the rate at which water is lost is often poor. This is bee other factors such as the number of stomata and their opening and ng influence water loss.

TURE OF FRUITS AND VEGETABLES

as well as on the occurrence of supporting tissues and the cohesiveof the cells. Turgor is that pressure of the cell contents on the parelastic wall of a cell, tending to produce rigidity. It is produced by a
are balance of forces which maintains the cell at a normal volume yet
vs the exchange of substances. When the cell volume diminishes, the
becomes soft and flaccid; but if the volume increases beyond the point
can be accommodated by the elasticity of the cell walls, the cell
ures, the cell contents flow out, and rigidity is lost. The substance
ly responsible for changes in volume is water. When a plant wilts,
r has been so extensively lost from the cells that they no longer have
hal turgor but are soft and flabby.

re of the best known forces affecting cell volume is osmosis. Plant cell are for the most part composed of cellulose permeable to many types sizes of molecules. Inside the cell protoplasm may be stretched around ze storage vacuole. The protoplasm and cell wall act as a semine ple membrane, allowing water and some other small molecules to ahrough it. Water diffuses in greater amounts from a region in which the concentration (dilute solution or pure water) to one in which it is low in concentration (a relatively concentrated solution). The ole contains soluble compounds as well as colloidal substances: and if

the intercellular fluid is composed only of water or of a dilute soluwater will move into the cell. While the fruit or vegetable is still pathe plant, a fine balance is achieved and the volume of the cell is a tained at a certain normal turgor. The effect of a concentrated solution cell turgor is readily demonstrated by placing cucumber slices in a centrated salt solution or sprinkling sugar on strawberries. Both soo come limp as water rapidly passes from the cells into the solution.

Cell turgor depends on a number of factors:

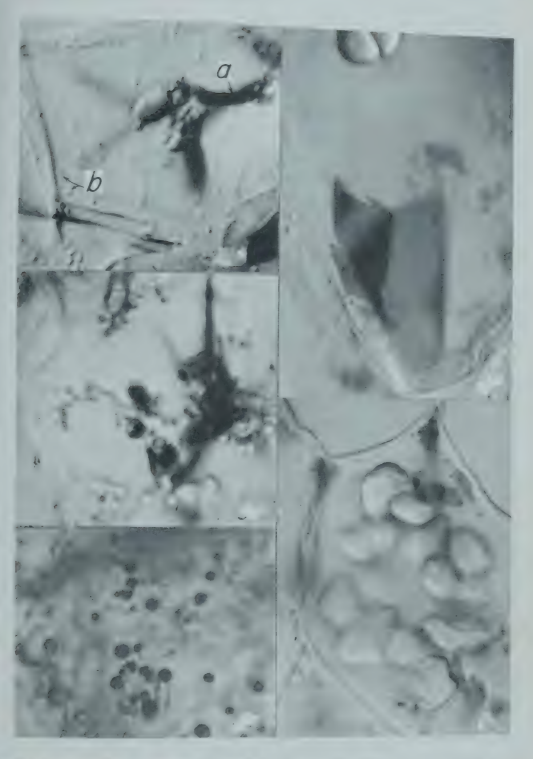
- (1) The concentration of osmotically active substances in the vacuo both true solution and colloidally dispersed.
 - (2) The permeability of the protoplasm.
- (3) Elasticity of the cell walls. If the walls are relatively high in elast a considerable increase in volume of the cell can occur before the cell tures. It will simply become more turgid as water flows into the cell. It ever, in the face of a shrinking volume a highly elastic cell wall will of a rapid loss of turgor for the tissues as a whole even as it adapts itsee the small volume of the cell. A stronger but more rigid wall will maintain firm texture even when the cell volume decreases.

When a fruit or vegetable is cooked, the protein is denatured, the die, and the vacuoles are no longer covered by a living membrane of toplasm. The protein usually precipitates and the permeability is mark affected. Often solutes and water stream out of the vacuoles into the incellular spaces or even out of the fruit or vegetable and the food becovery soft. However, when starch granules fill the vacuoles, they are slarge molecules that they cannot escape unless the cell walls are bro During cooking the starch granules swell, become gelatinized and remoisture; and if there are sufficient of them and if the process continutely may hold the water in the vacuole and maintain a firm texture. Hever cooked starch granules are never able to maintain crispness; the noticeable in a sweet potato and in peas and beans.

The rigidity of the structural tissues and of the cell walls is an portant factor in the texture of a fruit and vegetable. The changes in compounds which strengthen the structure of the plant with cooking be considered in the section on cooking changes. See p. 265.

PIGMENTS IN FRUITS AND VEGETABLES

The color of fruits and vegetables is exceedingly important to pleasure at the table. It is not only the subtle variety of flavors that m fruits and vegetables so pleasing, but also their variety of delicate bright colors.



PI 7.3. PIGMENTS IN PLASTIDS. Upper left: Chromoplasts a and oil droplets b in prienchyma cell from carrot root. Middle left: Destruction of chromoplasts upon a cell. Lower left: Carotene in solution in oil droplets in dead cell from carrot root. right I living carrot cell showing plate of carotene. Lower right: Living carrot cell in a starch grain with associated carotene plates.

Courtesy of Elliot T. Weier.

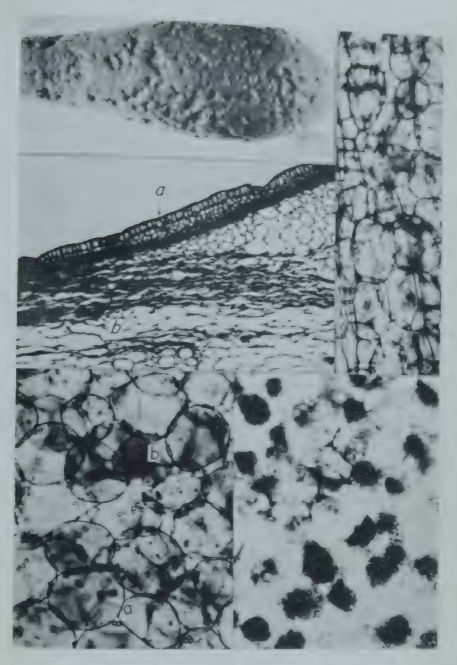
Most of the pigments occur in plastids, specialized bodies lying protoplasm of the cell. See Figure 7.3. For example, the chlorophylls in the chloroplasts, which under the microscope can be seen next cell wall as distinct bodies containing flecks of green pigment. Occasional pigment is present in the protoplasm as a crystal. Thus in carrot platelets of carotene can be observed and in tomatoes, needles or platelets of carotene can be water soluble pigments are dissolved in vacuoles, and not generally distributed through the cell.

In the early nineteenth century the only colored materials availabed dyeing were natural products. Much of the early research was direct an understanding of these natural dyes. When synthetic dyes were duced, interest in natural pigments declined, although it never disapped During the past thirty years there has been a tremendous interest in chemistry, and a great variety of problems have been studied vigored Among them is research in the pigments of plant and animal tist. The number of pigments whose structure and reactions have been pletely or partially elucidated is enormous. Only a relatively small ber of plant pigments will be considered here and only a very bried troduction to the field attempted. The number of studies on the chain pigments in foods is relatively small. But the results of the chemist be used to understand some color changes which occur when food cooked or processed.

The chief pigments of fruits and vegetables can be classified as (I carotenoids, (2) the chlorophylls, (3) the anthoxanthins, and (4) the arcyanins. Each group of pigments will be discussed and a brief introducto their structure, stability, and the changes which occur during cocconsidered. Tannins often account for the formation of off-colors deprocessing and they, too, will be considered.

The Carotenoids³⁴

The carotenoids are a group of yellow, orange, and orange-red fat ble pigments widely distributed in nature. In green leaves they occur i chloroplasts, small bodies close to the cell walls of the palisade cells. It cells are next to the epidermal cells on the upper side of the leaf. carotenoids are present in the lipid material along with the chloroph. The green color of the chlorophyll masks the yellow to red color of carotenes except in very young leaves while the amount of chloroph small. The bright, fresh yellow-green color of spring leaves is the resucarotenoids and small amounts of chlorophylls. These pigments are present in a wide variety of fruits—peaches, banana skins, tomatoes peppers, paprika, rose hips, squash, etc.—as well as other parts of plants.



124 7.4. Comparison of Living and Nonliving Plant Tissues and the Effects Upper left: Whole summer squash showing pitting injury due to 16-day storage (2°F) and one day at 21°C (70°F). Mann and Morris, 1947. Upper right: Parencells from cold storage summer squash, mounted in dilute neutral red. Cells are and therefore do not accumulate dye. Mann and Morris, 1947. Middle left: Showing through an injury pit of cold storage summer squash. a, epidermis, b, collapse of the forming the pits. Mann and Morris, 1947. Lower left: Living parenchyma cells, d r neutral red. Note accumulation of dye, b. Mann and Morris, 1947. Lower right: parenchyma cells, mounted in neutral red and strong sugar solution. Note plasmocalis. Mann and Morris, 1947.

Courtesy of Elliot T. Weier

in carrots, sweet potatoes, and in most yellow, orange, and red flow When they are consumed by animals, they tend to concentrate in and hence are found in blood, milk, egg yolk, and depot fat.

The name carotenoid is applied to all pigments chemically related carotenes, which were the first isolated. In 1831 Wackenroder extraor pigment from carrots and called the fraction carotene. We now know this "carotene" is a mixture of three isomers, α -, β -, and γ -carotene.

The carotenoids are either hydrocarbons or derivatives of hydroca and are composed of isoprene units. Isoprene is a diene,

and this molecule is the unit out of which the carotenoids are constructed to the carbon atoms while many, although not all, of the carbon atoms or eight isoprene units. Some of the cules contain a long unsaturated hydrocarbon chain with a ring at or both ends of the chain. Some of the molecules are symmetrical, so the folded in half the left half would be the mirror image of the right β -carotene and lycopene are examples of symmetrical molecules.

$$\beta$$
-carotene and lycopene are examples of symmetrical molecules.

 H_3C
 CH_3
 CH_3

Notice that β -carotene and lycopene differ only in the cyclization of end carbons of lycopene to form the rings of β -carotene. β -Caroten widely distributed in plant materials. It is readily prepared from car sorb apples, or paprika. It sometimes occurs free, although it is often companied by small amounts of α - and γ -carotene (see Table 7.2). It pene is the orange-red pigment of the tomato, but it is also found in hips, watermelon, apricot, and many other plants. It also is usually accompanied by other carotenoids.

Some carotenoids have two terminal rings or groups which are diffe Examples are α -carotene and γ -carotene.

ne difference between α - and β -carotene is in the placement of the ble bond in ring 2. Unlike these pigments γ -carotene has only one ring; of its molecule is like lycopene and half like β -carotene. Both α - and rotene occur in nature associated with β -carotene, although usually in ler amounts. In a very few plant products α -carotene predominates, example, in red palm oil it accounts for approximately 30 to 40 per of the carotenes present.

ber of xanthophylls have been isolated from plants and their structures mined partially or completely. They are often associated with caroice contain not only the hydrocarbon carotenes as their yellow tents but also the closely related xanthophylls. Cryptoxanthin is an exice of one of the xanthophylls.

ce that it is very similar to the carotenes and that it also is composed oprene units with the same system of conjugated (alternating) double 1s. Indeed, it differs from β -carotene only in the presence of the hyyl group. It often occurs as an ester of fatty acids, or it can be found tree form. It is one of the chief pigments of yellow corn, paprika, and the mandarin orange.

ere are also carotenoids which contain oxygens in other groups. Some etones, others are carboxylic acids, or hydroxycarboxylic acids. The ber of carbons is not always 40 as in the carotenes and xanthophylls

nay be smaller.

An example of a shorter chain pigment is crocetin which occur glycoside, crocin, in the spice, saffron. The deep orange-yellow cosaffron is not the result of crocetin alone since lycopene, β -carote carotene, and zeaxanthine are also present. Crocetin is:

O
$$CH_3$$
 CH_3 CH_3 CH_3 CH_3 CH_3 CH_3 CH_4 CH_5 CH_5 CH_6 CH_7 CH_8 CH_8

In crocin the carboxyl groups are esterified by carbohydrates (gentiob The number of carotenoids that occur in nature is probably very Along with those described, a number have been isolated and their tures either completely or partially elucidated. Information on carotenoids can be found in some advanced texts in organic chemistr in the original literature.

Aside from the beauty that plant pigments impart to fruits and tables, some of the carotenoids are important in nutrition as precurso the synthesis of vitamin A in the body. Vitamin A_i is identical with half of a β -carotene molecule plus a hydroxyl group, and in animals molecule of β -carotene is converted to two molecules of vitamin hydrolysis.

 β -Carotene

Vitamin A

the efficiency of the conversion has been the subject of extensive study. I used, and (3) the solvent for the carotene fed. In animals deficient in min A the conversion may be as great as 70 to 80 per cent; but if an mal has a large store of vitamin A in its tissues, if, in other words, it is then excreted in the feces. Rats show a great capacity for the conston of the carotenes to vitamin A, while dogs have only a small canty and cats none. If the carotene is fed in vegetable oil, it is readily orbed and converted to vitamin A; but if it is dissolved in mineral oil, it apes in the feces. In American diets the quantity of carotenoid presors is of great importance in total vitamin A nutrition.

Vitamin A2

The other carotenoids which are vitamin A precursors are those which e the same terminal ring as that in vitamin A_1 : α -carotene, γ -carotene, cryptoxanthin have one ring to each molecule. They are not as valuable

TABLE 7.2. DISTRIBUTION OF CAROTENOIDS IN SOME FOODS

of (Daucus Carota)	α -carotene, β -carotene, γ -carotene, xanthophyll, two hydrocarbons of unknown composition.	
at Germ	Xanthophyll, carotene (?).	
(Zea mays)	Zeaxanthin, cryptoxanthin, xanthophyll, α -carotene,	
-	β -carotene, γ -carotene, κ -carotene (?), neo-crypto- xanthin, hydroxy- α -carotene.	
cot (Prunus armeniaca)	β -carotene, γ -carotene, lycopene.	
h (Prunus persica)	β-carotene, cryptoxanthin, xanthophyll, zeaxanthin, unknown carotenoid.	
Bean (Glycine max)	α -carotene, β -carotene.	
pea (Vigna sinensis)	β-carotene, xanthophyll.	
ge (itrus aurantium)	β -carotene, lycopene, cryptoxanthin, xanthophyll. violaxanthin, zeaxanthin, β -citraurin, citraxanthin.	
afruit (Citrus grandis)	β -carotene, lycopene. α -carotene, β -carotene, xanthophyll, violaxanthin.	
penner or Chili	Capsanthin, α -carotene, β -carotene.	

ap icum fructescens)

as β -carotene in the synthesis of vitamin A, since they are capable of forming only one molecule of vitamin A for each molecule of pigment, but the are nevertheless significant. The other carotenoids do not contain this important ring and are therefore not precursors of vitamin A. The depth color of a fruit or vegetable has been suggested as a rough index of invitamin A value. But it can be seen that such an index could lead to gross errors since so many carotenoid pigments have no activity at all. For example, the yolk of the hen egg contains some vitamin A and is brightle colored because of the presence of two carotenoids, lutein and zeaxar thine, which are the dihydroxy analogs of α - and β -carotene, respectively. But these pigments have no vitamin A activity and the depth of the orang color of the yolk is no measure of its vitamin A value. Table 7.2 show the distribution in some foods.

The carotenoids are insoluble in water but soluble in lipids and in lipid solvents. In processing fruits and vegetables, loss of these pigments into cookery or canning water is very slight. They do undergo oxidation when exposed to air, so that in drying fruits or vegetables which contain these pigments a problem is sometimes encountered. For example, carrots and apricots show loss of pigment on drying. These pigments do not undergo hydrolysis except when they occur in plant tissues as esters, and they are not affected by changes in pH.

In blanched carrots, or those blanched and then dehydrated, the loss opigment is rapid. Carrots diced small and stored at 62° C in moist air afte blanching show a complete loss of pigment in 20 hours. Weier and Stocking have studied the change and find that antioxidants partially protect the pigment from deterioration in carrots ground to 48 to 16 mesh but have little effect on diced carrots. They were led to try antioxidants because carotene occurs in processed carrots (but not in the living cell) in fat drop lets; and they reasoned that the degradation of the pigment might be as sociated with oxidative changes in the fat. In leaves the enzyme lipoxidase is important.

Numerous other studies of the disappearance of carotenes have also been made. Usually the loss, if it occurs, is small—5 or 10 per cent. Altogethe the carotenes present few problems; they are bright and attractive and their color during food processing is easily maintained.

In a discussion of changes in fruits and vegetables on processing, one other aspect that should be mentioned is the distribution of the pigmen after processing. Weier and Stocking discuss the work of Weier and o Reeve on the carrot. The carotenoids in carrots are present in the chromo plasts. In some cells starch is present and the granules may more or less surround the carotene crystals. In the outer cells of the root the carotene

centration is highest, there is little or no starch, and the carotene is ent in crystals as needles, tubes, flakes, or spirals. These cells also continy fat droplets. When the cell is killed by blanching, drying, or chemreagents, the chromoplasts disintegrate and the carotene dissolves in oil siets. The oil droplets are sometimes originally present in the living cell may separate from the protoplasm after it is killed.

orophylls

ne green pigments of leaves and stems are usually held close to the cell in small bodies, the chloroplasts, along with some carotenes and hophylls. Two chlorophylls have been isolated, chlorophyll a and chloryll b; and they occur in plants in the ratio of approximately 3a:1b. mically, they are very similar. They belong to that group of important ogical pigments the porphyrins, which includes hemoglobin. They are y large molecules composed of four pyrrole rings held together by hene carbons (—CH=) to form a large flat molecule. In chlorophyll, a mesium atom is held by the nitrogen on two of the rings by ordinary contons. The other two nitrogens share two electrons with the magum to form a coordinate covalent bond (indicated by dotted lines). The nulas for chlorophyll a and chlorophyll b are given below:

$$CH_{3} = CH = CH_{2}$$

$$CH_{3} = CH = CH_{3}$$

$$CH_{3} = CH_{3} = CH_{3}$$

$$CH_{3} = CH_{2} = CH_{3}$$

$$CH_{3} = CH_{2} = CH_{3}$$

$$CH_{3} = CH = CH_{2}$$

$$CH_{3} = CH_{2} = CH_{3}$$

$$CH_{3} = CH_{3} = CH_{3}$$

$$CH_{3} =$$

You will notice that the chlorophyll a is an ester. The propionic residue position 7 is esterified with phytol, while the acid joined to position 6 esterified with methanol. Phytol, or phytyl alcohol, is an interesting himolecular weight alcohol because it occurs as an ester in a number of pla molecules. It has a chain which, although almost completely saturated, has a arrangement of carbons very similar to that in the carotenoids.

Phytol

Chlorophyll b differs in the occurrence of a —CHO (formyl) group in potion 3 in place of the methyl group (—CH₃) in chlorophyll a. Neith chlorophyll has been synthesized as yet.

The chlorophylls are of great importance in the plant because of the role in photosynthesis and the formation of carbohydrates from carbo dioxide and water. This role has been recognized for almost a hundre years and a tremendous amount of work has been done on the mechanism of photosynthesis. Only now are the reactions and the energy relations being elucidated.

The chlorophylls and other pigments are not the only components preent in the chloroplasts. One analysis of chloroplasts of leaves found 47 per cent lipid, 37.4 per cent protein, 7.8 per cent ash, leaving 7.1 per cent undetermined. A more detailed analysis of the chloroplasts of cabbase leaves reported 9.3 per cent chlorophyll, 0.5 per cent carotene, 0.8 per cent xanthophyll, 17.5 per cent glyceride fatty acids, 12.3 per cent wax, 4 per cent sterols, 13.3 per cent undetermined unsaponifiable, and 18.4 per cent calcium phosphatides. It is now believed that the chlorophylls exit in the chloroplasts as conjugated proteins.

The chlorophylls are very unstable molecules when the living cell killed and when the chemical and physicochemical relations in a cell a changed. Consequently, the chlorophylls are difficult to retain during ar food processing and special care must be taken to produce food that retain a bright, attractive green color. Some workers have suggested that the chlorophy is released into the cell. In 1940, Mackinney and Weast⁴⁵ reported that

ach and string beans which were carefully sectioned after cooking the roplasts were shrunken and often collapsed onto the protoplasm, but were still intact and the chlorophyll was still discernible in the plastids, chlorophylls exist as protein complexes in the living cell; and when the skilled by heating, the protein is probably denatured and the chlorophyll to the rapidity with which chlorophyll to in cooking food such as green beans. There is also the likelihood that permeability of the membrane surrounding the chloroplasts changes on leath of the cell.

hlorophyll changes to an olive green color and then to brown. The t likely reaction is the substitution of hydrogen for the magnesium has been complexed in the porphyrin to form pheophytin. It is uny that hydrolysis of either of the esters can be involved. Ester groups lly hydrolyze slowly and this reaction of chlorophyll occurs rapidly. The reaction is rapid in acid solutions and does not occur readily in sing waters where the pH is 8 or more. Esters, on the other hand, are e readily hydrolyzed in alkaline solutions than in acid. The possibility sidation as the reaction is likewise small since the brown color develops onditions where there is little free oxygen available. It is thought, afore, that when a vegetable becomes olive green on cooking, the chlosyll has formed pheophytin. The reaction can be written schematically ellows:

COOCH₃

$$+ 2H^{+} \rightarrow C_{32}H_{30}ON_{4}H_{2}$$

$$+ Mg^{++}$$

$$+ Mg^{++}$$

$$+ COOC_{20}H_{39}$$

ackinney and Weast studied this change in green beans and peas and npted to measure the amount of pheophytin produced in the food at ous lengths of time during the cooking process. Their results with tucky Wonder beans are presented in Table 7.3.

hen green beans are first dropped into boiling water, they, like other regretables, show a change in color. The green brightens, the velvety arance disappears, and the beans become more translucent. These tes are probably caused first by the wetting of the fine hairs on the of the bean. Washing the bean and rubbing it produce the same reto a slight degree. Then as the bean is warmed, air is expelled and intercellular spaces collapse or partially collapse. As cooking continues,

TABLE 7.3. KENTUCKY WONDER STRING BEANS HEATED IN WATER AT 100°C

Time (Minutes)	Pheophytin (Per Cent)	Color	
0	0	Green	
5	7	Green	
10	37.5	Green	
20	72.5	Brownish green	
30	86.5	Yellowish green	
60	100	Yellowish	

MacKinney, G. and Weast, C. A., "Color Changes in Green Vegetables," Ind. Eng. Chen 392-395 (1940).

plant acids are liberated; and because the chlorophyll is released for the protein complex or because the membrane around the chloroplasts comes more permeable or both, the acids react with the chlorophylls form pheophytin.

Blair and Ayres¹⁰ conducted an extensive study of methods for preseing the color in commercially canned peas. They found that on process the pH of the peas fell from 6.6 to 6.1. By a preliminary treatment whraised the pH and by maintaining it about 8, the color was preserved little of the chlorophyll was destroyed. We also know that when grivegetables such as spinach or cabbage, which produce considerable vola acid during the early part of cooking, are cooked in a pot with a cover, color very quickly changes to olive green and then to a dull brown. But the lid is left off so that acids escape during the early part of cooking, corretention is much better.

Since it is possible to maintain the green color in the presence of alk you might wonder why it is always recommended that baking soda, sodi bicarbonate, be omitted in cooking vegetables. The answer lies, of cour in the fact that in food preparation the maintenance of color is only one the important effects. With a high pH in the cooking or canning water, puticularly if the cation is sodium or potassium, cellulose hydrolyzes rapid and the texture of the vegetable becomes very soft and mushy. Some of vitamins, particularly ascorbic acid and thiamine, are very sensitive to he ing at high pH's; and in cooking water to which sodium bicarbonate been added, the rate of destruction of these vitamins is accelerated.

Blair and Ayres, 10 in an attempt to preserve the green color in campeas and still maintain a firm, but not tough, texture of the skins a cotyledon, used a 30 to 60 minute immersion of the peas in 2 per cent (0 M) sodium carbonate, a blanch in 0.005 M calcium hydroxide, and pressing in a sugar-salt brine which contained a suspension of 0.02 to 0.025

resium hydroxide. The pH of the peas by this process is maintained and 8, but the cationic balance produces the desirable balance between softening effect of the sodium ions and the toughening effect of the ium and magnesium ions. The calcium and magnesium ions tend to with pectic substances to form salts that toughen the cell walls.

me metal ions react with the chlorophylls to form compounds with ht green colors. Ferric, zinc, and cupric ions will replace the magum in chlorophyll. They will also react with the pheophytins which formed from the reactions of the chlorophylls with plant acids and

duce the corresponding complex.

udies by Strain on chlorophyll reactions have significance for changes ookery. He studied changes in green leaves boiled for 1 or 2 minutes, or en and then thawed. After fresh mallow leaves are boiled, there is a reion in the amounts of chlorophyll a and b and the appearance of chloryll a' and b', isomers of chlorophyll a and b. Leaves frozen and then ed yielded no other green pigments, but there was evidence that neithe oxidative nor the hydrolytic enzymes are inactivated. On drying eaves in air for 24 hr at 20°C chlorophyll a and b formed not only rophyll a' and b', but pheophytins as well. The enzymes are inactivated rying.

FLAVONOIDS

ne flavonoids are a group of compounds widely distributed in the plant dom. Every tissue studied so far has at least one of these compounds closely related one while most have many of them. They are water bie and are often present in the juices of plants. Chemically the

Flavone

noids contain two benzene rings with a three carbon bridge. In most hree carbon bridge is condensed through an oxygen into an interlate ring. The benzene rings hold hydroxyl groups.

he true flavonoids consist of the anthocyanins which are the red-blue-

purple pigments of plants; the anthoxanthins which are yellow; the chins; and the leucoanthocyanins. The last two groups of compound colorless but readily change to brownish pigments. They are probabled "food tannins."

Compounds related to the flavonoids are numerous and are also distributed in nature. They are as follows: (1) cinnamic acid and the complex caffeic and chlorogenic acids. Quinic acid, which forms par structure of chlorogenic acid, is a common fruit acid.

(2) Coumarins, which contain only one benzene ring and a condens ring,

HO
$$C = O$$

Umbelliferone, a coumarin

Hydroxy acids, such as gallic acid, tannic acid, and others.

he true flavonoids differ in the state of oxidation of the three carbon lege which separates the two benzene rings. The most common anthonidin, cyanidin, and the most common flavone, quercetin, differ only in oxidation of the three carbon bridge and have the same hydroxyl ups in the same positions on the benzene rings.

Anthocyanins

40st of the red, blue, and violet pigments that occur in flowers, fruits, other parts of plants belong to the group of pigments known as anthonins. These occur in plant cells as glycosides which are ethers of monohurides sometimes with one monosaccharide moiety and sometimes two. The color results from the structure of the anthocyanidin which ombined with the monosaccharides. The carbohydrates commonly ded to the anthocyanidins are glucose, galactose, rhamnose, and oconally a pentose. Most of the anthocyanins are soluble in water, and it also on boiling with fairly concentrated mineral acid, a condition which of encountered in food preparation, that the pigments are hydrolyzed orm the anthocyanidin and the carbohydrate.

Anthocyanin → Anthocyanidin + Monose

nly three types of anthocyanidins have been identified in plant tissues, ough a number of methyl derivatives of these three have been isolated. rgonidin, cyanidin, and delphinidin are wide spread in nature, with idin the most common. Often a plant tissue will have a number of ments composed of the same anthocyanidin but differing in the carbohymotety. Formulas for the chlorides are presented at the top of p. 242. compounds are formed when the corresponding anthocyanin is olyzed with hydrochloric acid. The methyl ethers which have been timed, peonidin, syringidin, petunidin, malvidin, and hirsutidin, are all atives of the hydroxyl groups on the benzene ring. In anthocyanins

Delphinidin Chloride

carbohydrate is always attached to the hydroxyl on carbon 3 and sionally one or more additional hydroxyls are etherified.

Some of the natural pigments occur as esters of organic acids; p-hydbenzoic acid, malonic acid, p-hydroxy cinnamic acid, and p-cumaric have been identified. These esterify either one of the hydroxyl group the anthocyanidin or one on the monosaccharide.

The Robinsons^{43,64,65} published a remarkable survey of the distribution of anthocyanins in plant tissues in 1938 and 1939. Much of the research this field has been carried out in their laboratory at Oxford. Their many contains a list of the pigments present in many garden flower autumn leaves, and in edible and ornamental fruits. The data for enfruits are included in Table 7.4.

The great variety of colors, hues, and tints that occur in nature and subtle shadings on the cheek of a fruit or in a blossom are the result number of factors.

(1) At low pH these pigments are red; the hues may be different, but are all reddish. Thus pelargonidin (it occurs in the plant as pelargonis some other derivative) is orange-red in acid solution while delphiniding bluish red. At high pH's the anthocyanins pass through a violet and blue color. Some turn green and then yellow at very high pH's (not countered in plants). Karrer and Jucker³⁴ point out that the pigment the red rose and the blue cornflower are the same cyanin. However, it red rose the pigment occurs as a salt of an acid while in the blue of flower it is present as metal salts. Cornflowers also contain a flav

blue at pH 11

genin, which is colorless but causes blueing of cyanin. (When an alkali has potassium hydroxide is added to an anthocyanidin, it is believed the following reaction occurs:

1) The concentration of the pigment alters the hue. Robinson reports that n an acid solution of synthetic delphinidin is poured on filter paper, a te solution gives a blue color while a concentrated solution gives red. intermediate concentration gives a purple color. When tannins are presthis is more pronounced.

3) In the cell sap the anthocyanins are frequently adsorbed on colloidal icles, probably polysaccharides. The pH is stabilized and the color innced by this association. In blue corn flowers cyanin is adsorbed and

pH stabilized at 4.9.

1) Frequently the anthocyanins occur as mixtures. As the composition of mixture is altered, the hue changes. Thus blue grapes contain not only osides of delphinidin but also of syringidin, the dimethyl ether of hinidin.

·) Sometimes plant cells will contain not only the anthocyanins as pigment wo some of the anthoxanthins which may be yellow and often yellow renoids.

·) Fannins are often associated with anthocyanins and alter the color.

he red color of beets results from a mixture of pigments. In 1948 noff and Aronoff¹ reported the fractionation of the pigments of beets

TABLE 7.4. ANTHOCYANINS PRESENT IN EDIBLE FRUITS*

Botanical Name	Common Name	Anthocya
Cornus mas Linn.	cornelian cherry	pel. 3-mono
Ficus caricus Linn.	fig	cyan. 3-mon
Fragaria vesca Linn.	wild strawberry	pel. 3-mono
F. virginiana Duch.	wild strawberry	pel. 3-mono
Morus nigra Linn.	black mulberry	cyan. 3-mon
Musa coccinea Andr.		pel. 3-mono
Oxycoccus macrocarpus Pers.	wild cranberry	peon. 3-mon
Pisum sativum Linn. var. (pods)	tall pea	cyan. 3-5 di,
Prunus avium Linn. var.	sweet cherry	cyan. 3-mon
P. communis Fritsh. var.	almond	cyan. 3-mon
Punica granatum Linn.	pomegranate	delph. di-
Sambucus nigra Linn.	European elderberry	cyan. 3-pent.
		gluc and mo
S. racemosa Linn.	red berried elder	cyan. 3-, 5-di
Solanum melongena	eggplant	delph. 3-pen
var. erculentum		delph. 3-bio
Vaccinium Myrtillus Linn.		malv. 3-mon
V. Vitis-Idaea Linn.	whortle berry	cyan. 3-mon
Vitis hederacea	grape	cyan.? 3 suga
V. quinquefolia	grape	malv. 3-mon
V. vinifera Linn.	vinifera grape	malv. 3-mon
V. labrusca	fox grape	malv. 3-mon
V. riparia		malv. 3-mon
V. aestivalis	summer or pigeon	malv. 3-mon
	grape	
V. heterophylla Thunb.		malv. 3-5 di

^{*}From Lawrence, W. J. C., Price, J. R., Robinson, G. M., and Robinson, R.

into eleven different fractions on a chromatographic column. One pig betanin, has been extensively studied and appears to be the glucosi an amino anthocyanidin. It also shows a change from red to blue a pH is raised.

In Cookery and Processing. Fruits and vegetables which contains anthocyanins, the red and violet foods, present a problem in cooker processing because of the great solubility in water of the pigments. is always a tendency for the pigment to leach out in the cooking or calculate or to run out in the juice. However, if the cell walls remain in the pigment is not lost. Thus frozen red raspberries show excellent tion of the pigment, but in canned berries the color gradually passes.

[†]Structure of anthocyanin shows anthocyanidin and carbohydrate.

Pel. = Pelargonidin (3,5,7,4' tetra hydroxy flavylium chloride)

Cyan. = Cyanidin (3,5,7,3',4') pentahydroxy flavylium chloride

Delph. = Delphinidin (3,5,7,3',4',5' hexahydroxy flavylium chloride)

Malv. = Malvidin (3,5,7,4' tetrahydroxy 3,5 dimethoxy flavylium chloride)

Peon. = Peonidin (3,5,7,4' tetrahydroxy 3' methoxy flavylium chloride)

the canning solution until the berries are practically colorless. When ets are cooked without removing the skins or even cutting off the root, for retention is much better than when they are peeled and cubed. When they are hot pressed, most of the color is removed in the juice; and if they fermented in wine making for a short time before the pressing, the exaction is almost complete.

The effect of changes of pH on the color of the anthocyanins is often ticed in food preparation, and occasionally presents a problem. Most its contain sufficient acid so that the pigment remains red or bluish red ough the cookery or processing. But if a small amount of the juice is ded to dish water containing soap or a detergent and with a pH, there-e, of 8 or 9, a blue or a greenish blue color forms. When vinegar is ded to beets in pickling, the color often reddens. When red cabbage is oked in soft water with a pH near 7, there is a bluing of the color.

If vinegar is added to blue cabbage, the color will change back to red. hen red cabbage is chopped, blue coloration often occurs along the edges t. This is apparently not a reaction with iron ions, although a possibility, ce red cabbage macerated with a Waring blender shows the same change

the surface exposed to air.

If red cabbage is cooked in hard water or, more important, if the water s been softened and has a pH of 8 or slightly higher, the cabbage bemes a grayish violet. If the cooking water is distinctly alkaline, usually m the addition of soda, NaHCO₃, the color becomes green. Most of the thocyanins will form a green color at high pH's and sometimes even low at higher pH. These are irreversible changes, but we seldom see such olor change in food preparation since we do not encounter such high

The anthocyanins form salts with metal ions, and have colors that dend not only on the particular anthocyanin but also on the metal ion. It is of the colors are grayish purple. The reaction is particularly important in canning and often in cookery. When tin cans (actually tin-coated n) are used for canning those fruits or vegetables containing anthomans, it is necessary to lacquer the inside of the can. Tin cans without a lacquer will cause a discoloration of the fruit touching the side of the lacquer will cause a discoloration of the fruit touching the side of the latin pie plate is used for preparing blueberry pie, the bottom of crust becomes a grayish blue color. This is particularly noticeable if has been scratched so that some of the iron is exposed. Small detended to fruit acids rapidly dissolve the rust, and the amount excition with the anthoeyanins is increased. If a cherry pie is cut with a calcino with the anthoeyanins is increased. If a cherry pie is cut with a calcino with the knife is allowed to remain in contact with the juice, the latin purple is observed. When aluminum salts are added to fruit

TABLE 7.5. SOME OF THE IMPORTANT PROPERTIES OF THE COMMON ANTHOCYANIDINS

	Pelargonidin	Cyanidin	Delphinidin	Peonidin 3' Methyl Cyanidin	Malvidin or Syringidin 3'-5' Dimethyl Delphinidin	Hirsutidir 7-3'-5' Tri Delphinid
Color of Aqueous Solution	Red	Violet red	Bluish red	Violet red	Violet red	Violet red
Solubility of Chloride in Water	Readily soluble	Only slightly soluble	Very soluble	Readily soluble	Slightly soluble	Slightly soluble
Ferric Chloride Reaction	Not definite	Intense blue	Intense blue	Faint— not definite	No reaction	No reaction
Behavior Toward Fehling's Solution	Reduces when warmed	Reduces in the cold	Reduces in the cold	Reduces in the cold	Reduces when boiled	Reduces when boiled
Color Change in Soda Solution	Violet then blue	Violet then blue	Violet then blue	Violet then greenish blue	Violet then blue	Violet then greenish blue
Behavior in Aqueous Solution	Color fades on standing	Color disap- pears on heating	Slow fading in the cold; when heated, rapid fading	Color disap- pears on heating	In very dilute solution color disappears when heated	In very dil solution co disappears heated

juices which contain the anthocyanins, a change in color occurs. But usually not as marked as with iron. In cooking fruits for jams and jellical aluminum vessels some of the change in color may be the result of thaction with aluminum ions. Iron vessels cause a marked alteration in color of the fruit and must be avoided. Reactions of a few anthocyan are summarized in Table 7.5.

Notice Approaches and Strawberries. The deterioration in the color of glass-page and strawberries and strawberries. The deterioration in the color of glass-page and strawberries and strawberries and strawberries and strawberries. The deterioration in the color of glass-page pears, peaches, plums, and grape juice is much more dependent on temperatures and oxygen content than light; ascorbic acid protects

rs. plums, and peaches against discoloration and the development of flavors, but it does not protect grape juice.

Anthoxanthins and Flavones

one of the most important groups of pigments in plants are the anthotnins and the flavones. They are yellow pigments usually dissolved in cell sap. The anthoxanthins are glycosides which on boiling with dilute i yield one or two molecules of monosaccharides and a flavone or a one derivative such as a flavonal, flavanonal, or isoflavone. The basic structure of the flavones is:

vanols have a hydroxyl group in position 3; flavanonols do not have a ble bond between carbons 2 and 3; and isoflavones have the phenyl up in position 3 instead of 2. Most flavones contain a number of hyxyl groups and some contain methoxyls. In the anthoxanthins one or re carbohydrates, usually monoses or dioses, etherify one or more hyxyls.

hese pigments occur dissolved in the cell sap and are usually pale ow or colorless but occasionally bright orange. Most bright yellow or age truits and vegetables are colored by carotenoids rather than anthomatical and flavones. Nevertheless the two latter are widely distributed are probably present in all white fruit and vegetables as well as those ared green with chlorophyll or red, blue or purple with anthocyanins. In the past twenty years there has been considerable work on the orac chemistry of these pigments. Recently interest has centered on the of these pigments in plant metabolism, in the relation of the genes and

inheritance to flavone synthesis, and in their possible medicinal value

The number of flavones and anthoxanthins isolated from plants is la Seshadri⁶⁸ in a review in 1951 listed 18 flavones, 27 flavonols, 5 flavano and 14 isoflavones. Many of these are from plants or parts of plants used as food. The number isolated from foods are relatively small many common foods have not been studied.

Examples of some of the flavones which occur in this group are:

Quercetin occurs in onion skins, tea, hops, horse chestnut, sumach, rose, and the bark of the American oak and many other tissues. Its a cosides are likewise widely distributed. A few in foods are the 3-galactos in Grimes Golden and Jonathan apples, and 3-glucoside in corn. Ruis 3-rutinose (disaccharide of rhamnose and glucose) quercetin and curs in many grains, in tomato stalks, in rue, elderberry blossoms, Califnia poppies, etc. One of the flavones of lemons and tangerines is also derivative of quercetin. Apigenin is present as a glucoside in parsley a is also one of the pigments of the yellow dahlia. Glycosides of apige also occur in field daisies, cosmos, and zinnias. Hesperitin occurs oranges and lemons as a 7-rhamnoside.

The anthoxanthins are yellow to orange in color and are dissolved the cell sap of flowers, stems, leaves, roots, and even wood. They water soluble pigments and differ in this from the carotenoids, the ot group of yellow pigments, which are lipid soluble. The anthoxanthins in small amounts in many fruits and vegetables which we ordinarily sider white or colorless. For many years a simple test has been used to onstrate the occurrence of flavones and anthoxanthins in foods, and in in all plant tissues. A drop of sodium hydroxide solution or ama vapor is applied to a slice or portion of the plant with the foron of a deep yellow, orange, or brown color. This color was supposed now the presence of flavones. However, many other common plant ponents give this color change and the results are not specific to flas and anthoxanthins. Some hydroxy acids are very widely distributed many of them as well as some proteins give the color change. Howwhether the change in color is caused by the occurrence of anthohins and flavones or tannins, or both, it is of some interest in cookery. hanges in Cooking or Processing. When a food is cooked in alkaline ers we often see the development of a vellow or cream color. Hard er will often have a pH as high as 8 and softened water which contains ICO, in place of Ca(HCO₃), will have a pH even higher. Potatoes sed in softened water often have a creamy color. The occurrence of color can be prevented or the color removed by adding a little cream of er. If the potatoes are cooked in chunks, bands of yellow are sometimes where the pigment is more concentrated in some cells. Rice also shows yellowing when cooked in softened water, and it too can be kept a r, bright white by adding cream of tartar to the cooking water. This t is most noticeable in onions, particularly yellow skinned, for the turns pale yellow and the cooking water is bright yellow when alkawater is used. Cauliflower and cabbage likewise sometimes show a yelng. Occasionally cauliflower turns a brown or pinkish brown color on .ing.69 Part of the color may be caused by flavones, but most is the It of reactions of ions with the tannins. Tea contains both the flavone, cetin, and tannins. The change in color has probably most frequently noticed when the acid of lemon juice causes a fading of the color. any of the flavones have definite physiological effects on man. Indeed, ctive principle of some old drugs, used long ago, has been shown to ther one or a mixture of flavones or anthoxanthins. One of the most esting is the mixture which has been called vitamin P or citrin. 1936, Szent-Györgyi and his collaborators showed that a factor other ascorbic acid was important for the maintenance of normal capillary capility. They named the factor vitamin P for permeability. In its see guinea pigs and even rats develop fragile capillaries through h protein passes and which rupture readily to produce hemorrhages. claimed that human patients with capillary fragility that does not and to ascorbic acid are likely to be deficient in vitamin P. The factor

is often more abundant in the rind of lemon and oranges than in the which is a rich source of ascorbic acid. A concentrate isolated lemon peel was named citrin, a term still used occasionally. Citrin per to be a mixture of flavones and anthoxanthins. It is composed of he din, eryodictin, their glucosides, and rutin which is the rhamnoglus of quercetin. The effect is difficult to estimate and there has been coing opinion of the role of the flavones and of ascorbic acid in the tenance of normal blood vessels. A number of derivatives of flavones been tested and some seem to be effective. Other flavones have been sto be diuretics, vasoconstrictors, heart stimulants, and cathartics.

Tannins

In prehistoric times it was discovered that some plant substance able to react with components in the skins of animals and "tan" The leather produced is much more durable than the dried ski ancient times tanning materials isolated from plants became object commerce. Knowledge about extracting these substances, handling and applying them to skins developed through trial and error. In mo cent times as science has been applied to an understanding of ar skills, there have been attempts to develop methods for detecting these stances. The tannins react with a number of ions and form dark of which have been used for inks; they are readily oxidized with permang and can be titrated. Both the ion tests and the permanganate test nonspecific.

As interest developed in the chemistry of foods, dark colors and a gent tastes were ascribed to tannins. Since it was difficult to separate compounds responsible for the darkening or astringency at that time, the nonspecific permanganate and ferric chloride tests were used. certain compounds in foods react, the term "tannin" was used for even when it was recognized that these compounds probably had litt no reaction in tanning leather.

Today the tannins of foods appear to be comprised of the catechin leucoanthocyanins, and some hydroxy acids. All of them give colors metal ions. Those which are ortho and para dihydroxy benzene de tives are readily oxidized by permanganate although the mono hydro meta dihydroxy derivatives are not. The substances that react with proteins in skins and bring about tanning are probably polymers of chins with intermediate molecular weights. The low molecular weight pounds in many fruits and vegetables are related to them.

Catechin and epicatechin are reduced derivatives of flavones. The isomers in which the ring and hydroxyl are probable *trans* in cat

in but they are probably closely related to the catechins.

Catechin

Leucoanthocyanin

hins and leucoanthocyanins are present in the tissues of those woody studied such as apples, peaches, grapes, almonds, and some pears, they are absent in herbaceous plants. They are present in cereals ugh the amounts vary.

and cacao have been extensively studied with the newer analytical ods such as chromatography. Tea contains a number of compounds of hin and epicatechin esterified with gallic acid. The most abundant are oyl epicatechin and 3-galloyl epigallocatechin. Epicatechin has been fied in both types of cacao beans, Forastero and Criollo beans.

ne of the hydroxy acids are found widely distributed in plants. Caffeic the most common phenolic compound in leaves and one of its esters.

chlorogenic acid, is present in many tissues. Phenyl caffeate is anothmon ester of caffeic acid.

If catechin is heated with dilute mineral acids, it forms an amored precipitate which is highly insoluble. This is called "tannin r"phlobaphene" and is believed to be a polymer of the tannin. It is put that the reaction is a general one for tannins or for one class of the Phlobaphene is reported by Stansbury, Field, and Guthrie⁷⁴ to of small amounts in the red skins (testa) of peanuts. They found a large cent) amount of catechol tannins as well as some other pigments.

Colors in Foods from Tannins. The tannins are readily dispersed water, some in cold, to form colloidal systems. When a fruit is pas apples in the preparation of cider or grapes in making juice of the tannins flow out in the juice. In extraction such as the brewing of coffee some of the tannins are extracted. The exceedingly astringer produced by boiling tea in water for some minutes is probably the of a thorough extraction of the tannins.

When the tea and coffee are brewed with hard water, a brown brown precipitate forms on the surface of the liquid; and as the brown cools, it appears throughout the liquid. Instead of a clear, spark fusion, the tea or coffee is distinctly muddy. In iced tea this

and noticeable, and with some waters the brew may be so full of preand the color is very noticeable. These transformations are believed
in and magnesium ions of the tannins in the tea and coffee with the
sleves are formed. Whether the precipitate is a simple calcium or
ty speak of the formation of tannates. The change in color which ocwhen lemon juice is stirred into black tea is also believed to be the

casionally Seven Minute frosting flavored with coffee turns green. viton investigated the development of the color and attributed it to rmation of iron tannate. She found that the color occurs only when ngs containing egg are used. No color appears if the frosting is with cream of tartar, lemon juice, vinegar, or brown sugar. She susdithat iron is introduced on the egg beater.

reenish gray discoloration occasionally occurs in chocolate ice cream as been accredited to iron tannates formed from specks of rust from e cream cans and tannins in the cocoa.

ren evaporated milk is allowed to stand in an opened can for some it imparts a grayish green color to coffee. Gould studied this reached found that it depended on the formation of rust on the can—the is proportional to the amount of iron in the milk. This is probably tion of the iron ions with the tannins of the coffee.

land¹⁴ investigated the darkening which sometimes occurs in the prom of maraschino and glace cherries. He believed that the discolorative result of a reaction between tannins and iron or copper ions, alhabe presented no direct evidence that tannins in the cherries are usible for the reaction. He also suggested that the tannin in the barwhich the cherries are stored might be important. He found that two parts per million of cupric ion causes darkening of the cherries 25 parts per million turns them black.

k spots on canned sweet potatoes is attributed to reaction of tannins ne iron ions formed from the walls of tin cans. A gray color in sugar our when iron ions from the crushers react with tannins.

od, and fullness of flavor to the food. The greatest difference in action between cider apples and culinary apples is in the tannin conhecider produced from those varieties with a relatively high tanning that body and a mellow astringency while that from culinary apples pid.

The presence of tannins in hops has been a subject of considerable and discussion among chemists associated with brewing. Some che find no evidence for tannins in hops although others get reactions a uted to them. It has been claimed that the tannins in wort and bee combined with protein and therefore do not give the ordinary tests for nins. In one report where the tannins and the place of their introdu into beer was detected by the color reaction with ferric chloride in presence of gum arabic and ammonium hydroxide, it was found that hops and malts contributed tannins. In 14 different samples of bee quantity of tannins varied from 25 to 55 parts per million.

The chemical composition of tea and the organoleptic quality has investigated. There is no consistent parallelism between the two but annins are the most reliable guide to quality. In tea the tannins are believed to give body and a pleasing astringency to the beverage. Too rannin can, of course, be very disagreeable and tea, which is boiled olowed to steep too long can be puckery and excessively astringent. Epichol, catechol gallate, and 5-hydroxy catechol have been isolated from tea.

In wine many compounds contribute to the quality, and tannins are group. Valaer⁸² says "A naturally high tannin content is desirable slight astringency, in the rougher red_types (for example Chianti) tannin being derived principally from the seeds and skins." The groups the removed from the stems before fermentation begins so that exive astringency is avoided. Also the wine cannot be allowed to remark contact with the seeds for too long during fermentation, or too much nin is extracted. A number of papers have appeared on methods of meaning tannins in wines and brandies.

THE BROWNING REACTION

when fruit and vegetable tissues are injured in any way or cut peeled during processing, a darkening of the tissues called the brow reaction sometimes occurs. (See previous discussion p. 108.) This reach has been extensively studied for a few fruits and vegetables but for me has had little research. Some browning reactions are enzymatic and occur in fresh living tissue or at least in tissues that still contain a enzymes. Thus when enzymes are denatured by heat or any other against reaction no longer occurs. Consequently, although a fresh peach turn brown after peeling, a canned peach will not. However, other denings may not be enzymatic. Thus when orange juice is concentrate often darkens with a deleterious effect not only on the appearance but

flavor. This browning reaction is nonenzymatic; it occurs at temres above those which denature most enzymes.

natic Browning

matic browning occurs in many tissues whenever they are injured. jury can be the result of bruising, cutting, freezing, or disease. That the injured fruit or vegetable which is exposed to air undergoes a darkening. Joslyn and Ponting32 published an excellent review of the of our knowledge about enzymatic browning in fruit in 1951. They out that as early as 1895 Lindet recognized that the change in color ing in freshly pressed cider is enzymatic. Since 1895 numerous studwe been made of the enzymatic browning reaction in fruits and vegebut little definitive information has as yet been gained. To begin the problem is extremely difficult and much of the work has been qualitative or suggestive. However, the techniques of enzyme chemhave developed in recent years. During the past ten years great s have been made in elucidating many enzyme reactions, and doubtciore long the browning reaction will be understood.

eral theories of enzymatic browning have been suggested, but as yet nas been established to the exclusion of all others. During the 1920's carried out extensive investigations of enzymatic browning in She did not isolate the compounds which undergo the changes but did out qualitative tests for their presence, concluding that browning s through the effect of an oxidase on a catechol compound to form a peroxide or hydrogen peroxide. The hydrogen peroxide then oxisome other compound, the chromogen, to form brown pigment. studied the enzymes in numerous fruits and found some evidence e presence of an oxidase and a catechol compound in the fruits that n readily (apple, apricot, banana, cherry, grape, peach, pear, and berry), but no evidence of these enzymes in the fruits that do not in readily (lemon, orange, lime, grapefruit, red currants, melon, pine-, and tomato).

ner early work seemed to indicate that the enzymes are peroxidases hat hydrogen peroxide reacts in the presence of peroxidase with some ound to form the brown pigment. However, the possibility that perses are the important enzyme systems in browning is now doubted. 12 and Joslyn³² believe that in apples peroxidase activity counts for or none of the browning. Other attempts to find hydrogen peroxide or instrate its activity have not been successful.

day most evidence suggests that oxidation of phenols or polyphenols

by enzymes is the principal reaction in enzymatic browning. The clature for the enzymes which cause oxidation of phenols or of poly is not standardized. They are called "phenol exidase," "polyphe dase," "phenolase," and "polyphenolase." Most have been stu crude extracts. They are very sensitive to decomposition at ordination peratures and are consequently difficult to isolate. Undoubtedly t many different enzymes that can catalyze the oxidation of phenols a derivatives by atmospheric oxygen. Not all catalyze the oxidation same phenols. Thus extracts of both the Royal apricot and Sphinx a are capable of catalyzing the oxidation of catechol and pyrogallo ever, although the apricot extract did not have any effect on glucinol, the avocado extract caused slow oxidation. An extract of will catalyze the oxidation only of o-dihydroxy phenols and apple catalyzes the oxidation of catechol and pyrogallol but not hydrod resorcinol, or phenol. This and other data suggest that the phenola ent in fruit tissues is not always identical and that the substrat affect may well be quite different in different plants.

The substrates, the compounds affected by the enzymes, have no isolated. Those tested which act as substrates for the phenolas plexes include catechol, tyrosine, 3,4 dihydroxyphenylalanine, chlorogenic, gallic, and protocatechuic acids, urushiol, phlorog hydroquinone and a number of anthocyanins and flavonoids. My these compounds are widely distributed in the plant kingdom and not their presence or absence in many plants is under study, precursors brown pigments in specific foods will soon be known.

The course of the reaction or reactions is not fully known. Ox absorbed, carbon dioxide often evolved, a quinone formed and the

ent is a polymer. It is evident with a number of substrates that one formation is not the only oxidative reaction. Catechol, for exercise with 2.4 atoms of oxygen during the sequence while its oxidato the quinone requires only 1 atom of oxygen.

me fruitful work in the control of enzymatic browning has been done. see of an antioxidant during processing has been used with some suc-Trius in preparing peaches for freezing, the addition of small amounts orbic acid not only prevents browning but also prevent loss of flavor. Teet of the ascorbic acid may be that of an antixoidant.

fruits susceptible to browning should be processed as quickly as pos-Heating destroys the enzymes responsible for the reaction; thus when is canned or made into jams or jellies, the browning reaction stops on as the fruit is heated sufficiently high to denature the enzyme. The temperature necessary varies with enzyme, rate of heat, pH, and factors. Deoxygenation and vacuum closing have also been used to tish oxidation.

the preparation of fruit for freezing, sugars and sugar solution have successfully used to prevent browning. The sugar solution coats the and prevents direct contact with atmospheric oxygen. If sugar is added fruit, it dissolves in fruit juices and forms a concentrated solution into the Concentrated sugar solutions inhibit or depress the activity of sudases, among them the phenolases, although at lower concentrate activity is sometimes enhanced. Some sugars other than sucrose been studied.

The effect of many salts and compounds such as sulfur dioxide, gen sulfide, hydrocyanic acid, and thiourea has been tested and t fect on oxidases studied. Halides and sulfites inhibit darkening of Joslyn and Ponting³² give the following list of compounds common to prevent browning and another list of patented inhibitors:

- (1) Commonly Used or Unpatented Chemicals: cystein, cystine, thione, sulfonamides, sulfurous acid and its salts, sodium sulfide, chloride, citric acid, hydrochloric acid, ascorbic acid.
- (2) Patented Chemical Inhibitors: (a) Sodium thiosulfate: Elion
- (b) thioamides such as thiocarbamide: Denny (1943); (c) chloride, ascorbic acid, and sodium bisulfite: Johnson and Gu (1949).

Hydroquinone, toluhydroquine, hydroquinone + lecithin, quinone + triethanolamine, diphenylamine, pyrocatechol, p-phenol, or resorcylaldehyde: Johnston, et al. (1943).

o-hydroxy aromatic oximes free of strongly acidic groups: Do et al. (1943).

Product of condensation of one mole of an alkali primary amin phatic carboxylate with at least one mole of an o-hydroxy substantic aldehyde. Preferred deactivators are salicyl derivatives dium glycinate, disodium glutamate, sodium tyrosinate and structures cysteinate: Downing and Pedersen (1944).

Thiosemicarbazide and its derivatives: Clarkson (1946).

In the home preparation of fruits, pineapple juice and lemon juice long been used to prevent browning. Pineapple juice has a relativel percentage of sulfhydryl compounds which are active antioxidants lemon juice contains both citric acid and ascorbic acid.

The pH has an important effect on the rapidity with which browning.

Curs. Acid dips are sometimes used to lower the pH and by this not delay or retard browning.

Many studies have been made on the darkening of potatoes whe are exposed to air by grinding, grating, or peeling as well as on the ening that sometimes occurs when they are cooked. The reaction up posure to air is doubtless enzymatic. Wallerstein, et al. 85 showed the tendency of white potato juice to pinking or graying increased on ing unless it was blanched at boiling-water temperatures for 50-6 Some evidence has accumulated to indicate that the pigment formed melanin and that the enzyme is tyrosinase. Some investigators have gested that the reaction which occurs on cooking likewise is enzymate is the result of the rupture of the cells with the release of tyrosinal

s improbable, however, that a reaction occurring at such high temures and even continuing after potatoes are cooked could be enzyther workers have presented evidence that the reaction in cooked loses is nonenzymatic. This will be discussed on p. 261.

enzymatic Browning

nenzymatic browning has been the subject of numerous investigations nevertheless, this reaction or these reactions are also only slightly stood. There have been many attempts to control the browning that s on storage of citrus fruit juices; in grape, strawberry, and raspjuices; and in dried apricots, peaches, pears, and apples. There have been some studies of the possible course of the browning reactions in But the number of factors which must be controlled is so large, and exture of compounds in a fruit is so complex, that again knowledge is sentary. The subject is well reviewed by Stadtman.

ree hypotheses have been suggested to explain nonenzymatic brownhere has not been sufficient work as yet to completely rule out any pothesis or to declare any other correct. (1) The browning reactat occurs between carbohydrates and amino acids results in the form of brown pigments. It is known as the Maillard reaction and is beby many to explain the browning found in processed fruit. (2) Asacid undergoes oxidation with the formation of a compound which ces brown pigment. (3) Carbohydrates or carbohydrate acids (asacid is a carbohydrate acid) decompose to furfuraldehyde or related bunds which then polymerize or react with nitrogen compounds to prown pigment.

temperature of storage, the amount of moisture, and the exposure stuit or fruit juice to oxygen either during processing or storage luential in the development of browning. For example, dried apricots had been treated with sulfur dioxide were canned and stored at diftemperatures. The samples stored at 46.1°C (115°F) darkened in 3 but those stored at room temperature (approximately 21.1°C or did not darken for three months and those stored at 0°C (32°F) darkening after six months. Similar data have been accumulated until orange juice, ground dehydrated apples, strawberry, currant, pherry juices. The data indicate that, in general, increase in temperature speeds up browning.

lies on the influence of moisture on browning indicate that there is effect in preventing discoloration, but the data are somewhat consince Stadtman, et al. showed that in dried apricots it is not

moisture alone, but oxygen uptake as well, that determines how darkening will take place, it is possible that some of the conflicting are caused by lack of control of available oxygen. Nevertheless, it a that lack of moisture favors the development of darkening.

Oxygen uptake either during processing or during storage has gene been shown to be a factor in browning. Some oxidation is apparent sential for the development of the reaction. For example, when the space in canned orange juice is increased, the rate of browning is portional to the increase in the space. Dried fruit is always expooxygen during processing.

There has been some attempt to discover what compound origoresent in the fruit is responsible for the browning reaction. The sibility that ascorbic acid is responsible has had considerable support the work done on citrus juices, particularly orange juice. The as acid content of the juice falls off as browning occurs. Addition of as acid to orange juice causes a marked increase in the rate of browning also been reported that a similar effect is shown in strawberry but other work indicates that grape, apple, and cranberry juice show creased browning rate when ascorbic acid is added.

The possibility of a reducing sugar as the reactant has been studing a number of workers. Slight decreases in these sugars appear to occur orange juice concentrates as browning occurs. However, attempts move the sugars by fermentation from orange juice had little influent the rate of darkening. Stadtman and his group were able to remove the ducing sugars in apricot syrups by fermentation but were only able to crease the rate of the browning reaction by about half. There is at put the possibility that both ascorbic acid and reducing sugars may be portant in the browning reaction.

The possibility that furfuraldehyde and its derivatives are involved the browning reaction in apricots was tested by Stadtman and co-wo They continuously extracted apricot syrup with ethyl acetate in furfuraldehyde is soluble, and during the extraction period there we browning. When extraction was discontinued, browning occurred. The tract contained a number of compounds among which were furfuraldeand hydroxyl methylfurfural.

Furfuraldehyde

Hydroxy Methylfurfural

n furfuraldehyde is added to apricot syrup, the rate of browning is

me work has been done on the isolation of the brown or black pigformed during browning. By repeated precipitation, a product has obtained from dried apricots which appears to be homogeneous. The nature of the compound is still unknown.

e black pigment formed when potatoes are cooked is most noticeable the stem end. Much research has been devoted to this problem too. been found that some varieties of potatoes are much more prone tkening than others. Northern grown potatoes are much more likely e this browning reaction and careful evaluation seems to indicate that eratures below 70°F at time of maturation favor the reaction. The ocnce of more extensive blackening at the stem end of the potato apto be caused by a higher pH at this end of the potato. The pH of oking water is also of great importance, with more extensive darkenceurring at high pH's.

present, then, it is known that a nonenzymatic browning reaction s when some fruits and vegetables are processed. We know that the on is influenced by atmospheric oxygen, moisture, and temperature. Impounds undergoing change may be ascorbic acid or some of the ing sugars. It is quite possible that this is not a single reaction, but up of reactions with one of them more important in certain fruits: ver more needs to be learned about the reaction. It is of consider-conomic importance to processors of these foods since not only is the trance of the product seriously affected but the flavor and the aszacid content as well.

IC SUBSTANCES

the substances (see discussion of structure, p. 87) are widely disced in plant tissues and are present in rather large amounts in rapidly ng succulent tissues with a high water content. Some cells possess composed of a single layer, while others on cell division form walls osed of three layers: a primary wall, a secondary wall, and a middle a that is shared with adjacent cells. The middle lamella is believed to nposed chiefly of pectic substances that act as a cementing material let the cells together. By staining tissue sections one can readily see walls and middle lamella under the microscope and the relative

walls and middle lamella under the microscope and the relative css or thinness can be easily determined. But when an attempt is to demonstrate the composition of these layers, the results are not so ut In many studies a stain, ruthenium red, has been used to reveal

the pectic substances. It has been shown, however, that although this does color most of the pectic substances, it will also react with other pounds. Therefore, the work with staining techniques is still open to tion since a stain that colors all the pectic substances yet does not other substances has not been found.

It is believed that the middle lamella is composed of insoluble substances, probably calcium pectates. The primary wall of many carich in pectic substances believed to be protopectin. These substances of chemical changes as the cells mature and the fruit or veget ripens. Many parenchyma cells contain large vacuoles filled with cell Some investigators believe that pectic substances are present in the ce and that this may even be the source of the pectic substances in the and middle lamella.

Apples have been extensively studied in all stages of growth and ma tion of the fruit, both by staining and by isolation of pectic substa Even when the fruit is extremely small, 1.3-1.5 cm in diameter, the m lamella and cell walls are readily stained with ruthenium red. This obs tion has been interpreted to indicate the presence of pectic substances i middle lamella and the cell walls. As the fruit grows, the total pectic stances increase; however after harvest as the apple softens, the amount soluble pectates and pectinates increase while the total pectic subst decrease. See p. 285. The crispness of the apple and its ability to puncture are the result of a number of factors. However, the presen pectic substances in the middle lamella and in the cell walls is believe be of great importance. When the pectic substances diminish in the m lamella, the cells gradually loosen and can be torn apart more rea When they diminish in the cell walls, the walls become thinner and readily punctured. As the apple continues to soften and grow mealy pectic substances that can be stained finally disappear completely.

It has also been shown that in bananas a marked change in the passive substances occurs during ripening. On storage the amount of soluble passive substances increases, while the total pectic substances decrease. In on riety of bananas, it was found that there is only a trace of water so pectic substances in the green bananas; but the amount increases a banana passes through the ripening stages.

Similar increases in the per cent of soluble pectic substances in m peaches as compared to immature peaches has been measured for four rieties. There is a corresponding softening of the fruit as it ripens the pectic substance becomes soluble.

Pears also show changes in the distribution and total amount of p substances as they ripen. The soluble pectic substances increase a softens; as these compounds dissolve, the juice that can be pressed acreases in viscosity.

e viscosity of fruit juices is proportional to the amount of pectic ances dispersed colloidally in the juice. When tomato juice is pressed e cold-break method, the cells are macerated and enzymes released of the enzymes present in tomatoes are then free to catalyze the hysis of pectic substances. The first, pectin methyl esterase, catalyzes moval of methyl groups from the carboxyls on the galacturonic acid tes of pectinic acids with the formation of low-ester pectinic acids ztic acids.

Pectinic acids + H₂O pectin methyl esterase CH₃OH + pectic acid

esecond enzyme, depolymerase, then attacks the low-ester pectinic or pectic acid and by hydrolysis forms products whose molecular ts are too low to contribute to the viscosity of the dispersion. The break method forms a watery tomato juice. If the tomatoes are heated 180°F (82.2°C), the enzymes are denatured, and the pectic subservated with the juice are pectinic acids with a sufficiently high ular weight so that they form viscous colloidal dispersions.

entribute considerably to the ability of the juice to retain sediments pension. In tomato juice a desirable product has considerable susd materials and a viscous juice can hold these without their settling other factors such as the size and nature of the particles are imput, but juices with a watery serum do not have the ability to hold ather fine material in suspension.

ce a clear liquid is expected in some juices, the presence of a relable high per cent of pectic substances may be objectionable. If it is sary to filter the juice, a high viscosity slows down the filtration and resent an economic problem. The pectic substances are ordinarily reliable by the use of pectinases, enzymes which depolymerize the pectic notes until the molecules are so small they no longer form coldispersions or act as a protective colloid. "Pectinase" is an old term commercial enzyme preparation that contains a polygalacturonase her enzymes. The polygalacturonase hydrolyzes pectic acids to low har weight polygalacturonides and galacturonic acid. In prepared the pipe juice, removal of pectic substances is almost always necessary of the high percentage present. In wine the clarification and present of tartrates and other sediments is sometimes difficult because the substances hold them suspended. When the wine is treated with ase, precipitation occurs.

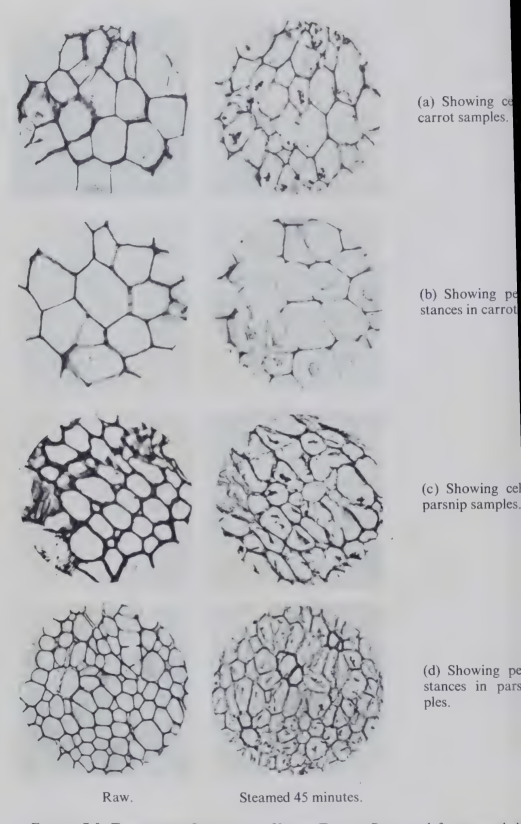


FIGURE 7.5. TRANSVERSE SECTIONS OF XYLEM TISSUE. Raw on left, steamed 4 on right. Magnification × 300. Courtesy of Simpson, J. I. and Halliday, E. G., search, 6, 189–206 (1941).

HANGES ON COOKING AND PROCESSING

Ine chemical changes that occur when fruits and vegetables are boiled tater, steamed, canned, dried, or frozen are still for the most part unges in the appearance, texture, flavor, and color of the fruit or vegete occur. But exactly what these reactions are is still, for the most to the work of the future. There have been a number of studies on popes and apples and a few on some other fruits and vegetables. The nges observed in a limited number of vegetables and fruits are sugtive of the reactions in others.

If As a result of changes in cellular structure, all fruits and vegetables ergo softening when cooked, no matter by what method. We can expect er that the cells are separating, rupturing, shrinking, or that a comation of these changes occurs. When cells separate during processing, cementing substances between the cells, in the middle lamella, must r. Thus, since the principal compounds in the middle lamella are beed to be the pectic substances, their change is the first one considered. since the cell walls are known to be composed of a matrix of cellulose erimes encrusted with other materials, the possibility of changes in ulose must be considered. (3) Starch granules can be seen in microscopic ions of most fruit and vegetable tissues and are known to swell rapidly vater. (4) The intercellular air changes in volume when subjected to her temperatures because of expansion. (5) Alterations in pigments, nation of acids, and release of low molecular weight sulfur compounds indicated from observations of gross changes. A large number of reacis ire suggested by cursory observation, but careful and conclusive inigation of their role is difficult and has been applied to relatively few ts and vegetables.

anges in Pectic Substances

and parsnips. See Figure 7.5. They determined the amount of pectic stances in the raw vegetable, vegetables steamed 20 minutes, and vegetables steamed 45 minutes and found that while there is a steady increase ne amounts of pectins and pectates as steaming progresses, there is a second in the protopectin as well as the total pectic substances. When this of tissue were imbedded in paraffin and stained to show the middle at vegetables steamed 45 minutes had a much thinner middle lamella the fresh tissue. Simpson and Halliday concluded that at least one are in the softening of carrots and parsnips that occurs on steaming is

the changes in pectic substances—the hydrolysis of protopectin to pect See Table 7.6.

Many researchers have attempted to explain the difference in the culir properties of some potatoes. Mealiness, waxiness, and sloughing are result of cell separating. It has been found that the addition of calc salts to potatoes during boiling greatly reduced splitting and slough However, Freeman and Ritchie¹⁷ did not find a correlation between amount of pectic substances in raw and cooked potatoes and their meness.

The hardness of the water used in canning beans may have a proform effect on their texture. Addition of 1,000 ppm CaCl₂ to water produce beans so hard and tough as to be unacceptable. It is believed that toughening action of calcium salts is through their effect on the persubstances; calcium pectates are insoluble.

The pectic substances are also of considerable importance in the fing of canned tomatoes, apples, and other fruits by calcium salts. Can tomatoes which are graded A and which therefore bring the highest p must be relatively firm, yet full-colored. However, during ripening, matoes pass through this period rapidly, soften, and on canning unde considerable maceration and shredding. The addition of small amounts calcium salts to the pack increases the firmness of the fruit. Calci salts can be added to the dip or placed in the can with the salt. Calci chloride is permitted in the United States at a level of 0.07 per cand salts such as calcium citrate, sulfate, or phosphate may be used equivalent levels calculated on the basis of calcium ion. This level has effect on the flavor of the fruit but produces a marked effect on the finess. The method is also used widely for firming both canned and from sliced apples as well as baked apples. It has also been shown to be effect

TABLE 7.6. EFFECT OF STEAMING ON PECTIC SUBSTANCES
OF CARROTS AND PARSNIPS*

Fraction of Pectic Substance	Raw		Steamed 20 minutes		Steamed 45 mi	
	Carrots (%)	Parsnips (%)	Carrots (%)	Parsnips (%)	Carrots (%)	Par (
Pectin	3.7	4.7	.6.0	6.1	8.8	
Protopectin	14.1	10.2	9.0	7.7	3.6	
Pectic Acid or Pectates	0.8	1.6	1.0	2.0	1.3	
Total Pectic Substance	18.6	16.4	16.1	. 15.8	13.7	1

^{*}Results represent the per cent of dry weight and are computed from the average of five weights of cium pectate in each case. From Simpson, J. I. and Halliday, E. G., "Chemical and Histological Studithe Disintegration of Cell Membrane Materials in Vegetables during Cooking," *Food Research*, 6 206 (1941).

ny other fruit products and will perhaps become commercially imint in the future. Raspberries are firmer if they are treated with calcium
before canning. The results are also good with canned potatoes,
ites, and olives. Kertesz, Tolman, Loconti, and Ruyle studied this ren extensively and concluded that the calcium ions react with pectic
and low-ester pectinic acids which have numerous free carboxyl
ps to form insoluble calcium pectate. It is interesting that mealy apples
nich the amounts of pectic substances are low do not show an increase
mness when they are treated with calcium salts.

eve and Leinbach⁶² have studied the saucing properties of apples and pectic substances but have not found a correlation between them. The uble protopectins were determined by the usual Carré-Haynes method, ensteins sauce rapidly and those from the 1948 crop disintegrated rapidly than either the ripe or green 1947 Gravensteins. Newtown ins and Winesaps resist saucing, while Delicious and Jonathans are inediate. Apples tending to sauce readily underwent complete cell sepanwhen they were vacuum infiltered with water. When a CaCl₂ some was substituted for the water, there was no increase in firmness, ever, in Winesaps, Newtown Pippins, Jonathans, and some Delicious ples there was appreciable firming. These results indicate that, algh the difference between saucing and nonsaucing apples is not enin the pectic substances, the firmness of some varieties is probably ndent to some extent on these compounds and their distribution. See e 7.7.

ou will remember that it has been shown that changes in pectic sub-

TABLE 7.7. SAUCING QUALITY AND PECTIC SUBSTANCES
IN APPLES*

and Storage Time at 33°F;	Cold Water Soluble Pectin (Per Cent)	Insoluble Pectin (Per Cent)	Saucing
ous. C. Grade (1947) 1 Month	0.11	().29	Intermediate
	0.09	().34	Intermediate
ous, Fancy (1947) 1 Month	0.09	0.16	Good
istein (1947) 10 Days	0.20	().3()	Good
istein. Green (1947) 16 Days	0.06	0.35	Good
istem. Green (1948) 7 Weeks		0.31	Intermediate
ar & Grade (1947) I Month	0.05	().3()	Intermediate
r Fincy (1947) 1 Month	().()6	0.42	Poor
Fippin (1947) 1 Month	0.12	().34	Poor
q: Lancy (1947) 1 Month	0.05		Роог
1. Lancy (1948) 1 Month	0.04	0.35	. Apples I Com

m 1 = R M and Leinbach, L R, "Histological Investigation of Texture in Apples, L Com a Influence of Heat on Structure," Lood Research, 18, 592-603 (1983)

stances occur as apples ripen and mellow. When apples are held in stafollowing harvest, there is a gradual increase in soluble pectic substate and a decrease in insoluble but the total remains fairly constant. As apples become mealy and over-mature, there is a decrease in the pectic substances and a change of the soluble to nonpectic compout (See p. 285). However, Reeve and Leinbach did find that the percentage cold water extractable pectic substances from Cortlands (a saucing applications) four times (0.31 per cent) what it is from Delicious (an intermediate—per cent).

Reeve⁶⁰ has also studied the seed coat of young peas. He finds agents that react with the pectic substances have an effect on the young peas but not on the mature. Peas grown in soils with an abundance calcium have tough coats. He found that as the pod becomes mature pectic substances of the middle lamella in the cells that make up seed coat become encrusted with pentosans or hemicelluloses. It is litthat in the mature peas, changes in pectic substances are not as real available for reaction, since the pectic substances are partially cover with hemicelluloses.

It therefore appears likely that the softness that occurs on cooking processing fruits and vegetables is partially the result of changes in pectic substances. As our teeth sink into a piece of fruit or vegetate few cells are ruptured; instead our teeth separate the cells. It is litthat the ease of biting and chewing results from the changes in the stances that cement the cells together, the pectic substances. The lamble molecules of insoluble protopectins are in some fashion hydrolyzed smaller, so-called soluble pectic substances which are able to form loidal dispersions in water. Reeve's findings on mature seed coats do disagree with this conclusion, since it is a frequent observation that woody vegetables cannot be softened by cooking.

There is the possibility, at present not established, that the pectic stances in other vegetables become encrusted with compounds such as hemicelluloses incapable of partial hydrolysis under the mild condition cooking. Scott⁶⁷ found that leaves, stem, fruit, and flowers of a numbe plants (not foods) contain a thin layer of material which she ca "suberin," lining the entire system of intercellular air spaces as well on the inner surface of cell walls. This substance or mixture is insolutin 80 per cent sulfuric acid and also in chromic acid and increases amount as the plant ages. Scott considered the material waxy in nature, offered no evidence for this conclusion. Whatever its nature, it is likely to compounds resistant to 80 per cent sulfuric acid are not hydrolyzed during and that this layer may account for the lack of softening to

etimes occurs. Until more evidence accumulates, we conclude that ges in pectic substances are among the important reactions that occur occssing vegetables and fruits.

nges in Cellulose

It attention has been directed to the possibility that the cellulose il walls undergoes some changes during cooking or other food-process-nethods. The resistance of the cellulose from such well-known sources atton fibers, to the action of hot water, has thrown doubt on the postty of hydrolysis. However, the work of Simpson and Halliday with ots and parsnips shows that on steaming, the cell wall thins out; they me that this indicates a change in cellulose.

nges in Starch Granules

though starch granules occur in most plant tissues in storage leucos surrounded by thin strands of cytoplasm, starch also occurs in the roplasts of leaf cells when light falls on the leaf. Chloroplast starch ought to be only transient; i.e., rapidly hydrolyzed, carried in son to the storage cells and then resynthesized into the storage starch, ng processing the starch granules undergo the same changes that occur starch and water are heated. See p. 105 for a discussion of these ges in simplified systems. During heating the starch granules swell if tient water is present. Tissue slices of steamed or boiled potatoes show len and often gelatinized starch granules. Occasionally when the cell is ed with iodine, the outline of a starch granule can be seen, but more the whole cell is filled with the gelatinized starch. Sometimes the rupture and the gelatinized starch streams out.

number of papers have been published on the changes in potatoes, cularly dehydrated potatoes. Weier and Stocking⁸⁹ studied (1) freshly ned potatoes, (2) dehydrated mashed potatoes of poor quality, and (3) drated potatoes of acceptable quality. They found that in all three (numerous samples were used) gelatinization of starch occurred alghoutlines of the swollen granules could be detected in most of the In freshly mashed potatoes most of the cells remained intact with gelatinized starch between the cells. However, in poor-quality pomany of the cells ruptured, resulting in much gelatinized starch benthe cells. When water is added to these dehydrated potatoes, the h granules swell, rupture the cells, and stream out between them and a thick, sticky starch paste. Weier and Stocking conclude that



FIGURE 7.6. STARCH GRANULES IN POTATOES. Showing G-gelled starch escaped frotured cells. Magnification \times 40 and \times 100. Upper left and right: Unstained, separate of boiled, mature Russet Burbank potatoes. Lower left and right: Unstained separate of boiled, mature White Rose potatoes. Note more cell rupturing in White Rose, starch can be seen in all unruptured cells. *Courtesy of R. M. Reeve.*

with little starch in between.

ceve* found that when White Rose (nonmealy) and Russet Burbank v) potatoes are steamed or boiled, it is difficult to demonstrate the nce of individual starch granules because they are so swollen. See re 7.6. He found that in the nonmealy White Rose potato cell rupturas more pronounced and there was much more gelatinized starch ben the cells. He concluded that the escape of gelatinized starch from ired cells causes a sticky or gummy texture in cooked potatoes. Howhe found that factors other than starch are also important. Young et Burbanks produced a texture in the steamed slices between mealy waxy, but sometimes sticky, and yet showed no escaped starch. He had ations that the starch in the two types of potatoes had different hydraproperties because of the difference in the colors obtained with iodine ing and the difference in the ability of some starch to escape without eable rupturing of the cells. In some specially treated samples of non-White Rose potatoes, starch appeared between the cells although or no cell rupturing occurred. This is probably low molecular weight ose. See Tables 7.8 and 7.9.

ne swelling of the starch granules in cells is often a factor in causing to break apart. However, the changes that occur in starch granules ig processing is the result not only of gelatinization of starch but also ity of amylases and hydrolysis of starch to dextrins and maltose. Mann Neier46 reported that starch of carrots, although variable in amount ocation, is generally found in the cells of the cambium. They observed rate of heating during blanching has a marked effect: thus, if rapid, ring less than 60 sec to reach 75°C, sections of carrot stain blue, with e showing the presence of starch; whereas if slow, sections stain le. red, or pink, with iodine indicating presence of dextrins. The differlies in the different temperatures at which carrot starch swells and the erature at which the amylase is inactivated. Although carrot starch has atmization temperature between 40°C and 50°C. Mann and Weier that the inactivation temperature of carrot amylase is about 75° C. . it the carrot is rapidly heated, the enzyme is inactivated before any sive hydrolysis occurs. On the other hand, if the carrot is heated v ne starch granules swell and considerable amylase activity is posbefore the temperature of inactivation is reached. Although the releof these phenomena to other fruits and vegetables is not really n. this work on carrots may be applicable; indeed in other fruits and able blanching is usually carried to a point where enzyme mactivaici urs.

TABLE 7.8. COMPARISONS OF CELL SEPARATION, CELL RUPTURING, AND STAINING OF STARCH IN HEATED SLICES OF POTATO TUBERS AND THEIR RELATION TO FINAL TEXTURE QUALITIES*

			THE IEALONE GOALINES	IL TEATORE GUALITIES	
Treatments and Raw Materials	Cell Separa- tion	Cell Ruptur-	Iodine Gelled	Iodine Color* External Solution or Filtrates	Texture of Tissue after
Roiled		۵	Staten	(Amyloid Blue)	Final Heating
Mature White Rose	slough in water	+ + + + +	reddish to blue-purple	+ + +	slightly mealy
Young White Rose	sloùgh in water	+ + + +	reddish to blue-purple	+ + + +	soggy soggy or sticky
Mature Russet Burbank	slough in water	+++	blue-black	faint, if any	mealy when
				`	severe
Young Russet Burbank	slough in water	++++	blue-black	+	slightly mealy to
Steamed					S
Mature White Rose	readily obtained on microslide	+++	reddish purple	+ + +	slightly mealy
Young White Rose	readily obtained on microslide	+++	reddish to blue-purple	+ + + +	generally sticky or gummy, some-
Mature Russet	Slight slonghing	-	3		* times waxy
Burbank	a	+	blue-black	generally	mealy
Young Russet Burbank	readily obtained on microslide	+	blue-black or dark purple	faint	slightly mealy to

"I can Reve R M "Histological Survey of Conditions Influencing Taxona in Processing France in Processing Processing Processing Conditions and Conditions and

rubbery, slightly sticky when slice is broken	rubbery, shees sticky to gummy when broken	leathery, granu- lar when slice is broken	rubbery, granu- lar to waxy when slice is broken		no final heating	no final heating	no final beating	no final heating
+ + + + +	+++++	faint, if any	faint		+	++	nene	none
blue-purple to black	reddish purple	blue-black	blue-black		blue-purple	blue-purple	blue-black	blue-black
Slight	slight to none	none	none .		none	none	none	none
readily obtained on microslide	readily obtained on microslide	readily obtained on microslide	readily obtained on microslide		slight, by stirring	slight, by stirring	slight, by stirring	slight, by stirring
Mature White Rose	Young White Rose	Mature Russet Burbank	Young Russet Burbank	Soaked 1 hr at 80°C in 0.5°°. Solutions of Pectic Solvents	Mature White Rose	Young White Rose	Mature Russet Burbank	Young White Burbank

Boiled after Soak-ing at 75°C I hr

Changes in Intercellular Air

The parenchyma tissue of fruits and vegetables is composed separated by tiny pockets and passages of air. In some fruits and tables the amount of air may be appreciable, in others quite small peaches may have as much as 15 per cent air while plums have ver When the fruit or vegetable undergoes processing, changes occur intercellular air. If the product is simply heated, the air will swel the cells apart, and often cause cracks in the food. Thus in a baked or baked apple, swelling occurs; however, on cooling, the product and cracks in the body are noticeable. Further changes may take p the cell walls become more permeable. Cell sap, the solution held vacuoles, may escape into the intercellular spaces, causing a change appearance and juiciness of the food. Thus the chalky appearance product changes to one of translucence as the air is replaced with war

This can be readily seen when green beans are placed in hot wa soon as the bean is submerged, it begins to look greener because of charge of air from around the hairs on the surface and perhaps fr tween some of the surface cells. In many foods juiciness is marked fected because the fluid that now fills the intercellular spaces escapes from the tissues. Weier and Stocking⁸⁷ point out that if the sap is insufficient to fill the intercellular spaces the chalky appearance not disappear; thus in steam blanching of cabbage it is retained. On a fruit or vegetable a third change may occur: the discharge of interair and the filling of the spaces with the cooking water. Reeve and bach⁶² were able to cause complete cell separation in saucing app vacuum infiltration of the apple tissue with cold water. They sim placed the air in the intercellular spaces with cold water, and in varieties of apples this caused the apples to disintegrate. Reeve⁵⁹ poi that the large intercellular spaces of the apple provide a tissue readily impregnated with syrups. He did not find that the amount o cellular air (20-22 per cent for Delicious, Newtown Pippin, and W 23-24 per cent for Rome Beauty; slightly over 25 per cent for Grave explained the difference in texture of these varieties on cooking in w steam.

Production of Volatile Acids

Many green vegetables contain volatile acids that are partially giduring cooking. These acids are of importance during cooking because have a marked effect on the color and flavor of the cooked vegetablid is placed on the cooking vessel, these acids dissolve in the steam

consess on the lid, drop back into the cooking water, and lower its pH. prophyll, which is very sensitive to any pH below 7, will change to ophytin and other olive green pigments. Green beans, spinach, or broccooked in a covered kettle will be browner in color than another ple cooked without a lid for the same length of time and under the conditions. When the lid is left off the vessel, these volatile acids are ally evaporated. They affect the flavor in two ways: (1) they have sourand flavor themselves, (2) they speed up hydrolysis of sulfur-containally cosides and produce distasteful sulfur compounds. The volume of the in which one of these green vegetables is cooked is likewise intents ince these volatile acids are readily soluble in water and can be seed by using generous amounts of cooking water.

he chemical nature of the volatile acids evolved during cooking is at sent unknown. Presumably they are low molecular weight organic acids as formic, acetic, propionic acid, and perhaps lactic acid. The amount cid evolved during the cooking of the vegetables has been measured andensing the steam in a regular distillation outfit and titrating the

evolved.

are for the most part unknown. They are probably dissolved in the sap since biting or squeezing a fruit or vegetable releases juice in which are present. However, in some fruits at least part of the acids must it in a different section of the cell from the anthocyanins that are discident the cell sap. When plums, blackberries, or blueberries are cooked, sigments redden. This indicates that the pH has dropped. Blackberries blackberry cobbler may become quite red. In some way acid and pigt which have been separated in the tissue have now reacted. When tables are cooked in water, the acids are quickly leached out.

est investigation of organic acids in fruits and vegetables has been red out on the whole fruit by macerating or grinding the tissue and exing the acids. They have then been identified by typical reactions or stonally isolated and identified. Malic and citric acid are the most comand are present in small quantities even in fruits and vegetables not the considered "acid." They appear to be present in all plant tissues. example, beets contain 0.11 per cent citric acid, while eucumbers have per cent malic acid and pumpkin 0.15 per cent. Oxalic acid occurs in a per cent malic acid and pumpkin 0.15 per cent. Oxalic acid occurs in a me leaves such as spinach, beet greens, lambsquarters, and purslane, of tartaric acid have been reported in fruits and vegetables as different acid have been reported in fruits and vegetables as different acid have been reported in fruits and vegetables as different acid have been reported in fruits and vegetables as different acid, while cucumbers have a per cent acid have been reported in fruits and vegetables as different acid, while cucumbers have a per cent acid have been reported in fruits and vegetables as different acid, while cucumbers have a per cent acid occurs in a per cent acid occurs

quantities of the acids vary with season, ripeness, and variety, of See Table 7.14, p. 287. Succinic, lactic, benzoic, and isocitric have als found in small amounts in numerous fruits and vegetables. Cran contain quinic acid that also occurs in many plant tissues as part molecule of chlorogenic acid. See Table 7.10.

A few acids which are volatile have been identified along with nur other types of organic compounds important in the flavoring of fru vegetables. Kirchner³⁸ lists both acetic and n-caproic acids in the et

TABLE 7.10. ORGANIC ACID CONSTITUENTS OF FOODS*

Food Items	Citric Acid (Per Cent)	Malic Acid (Per Cent)	Other A
Apples:			
Crab	0.03	1.02	_
Delicious	_	0.27	-
Grimes' Golden		0.72	-
Jonathan		0.75	_
McIntosh	·	0.72	_
Rome Beauty .	_	0.78	_
Winesap	trace	0.50	_
Yellow Transparent	0.02	0.97	_
Apricots, Canned	1.06	0.33	_
Dried	0.35	0.81	Trace of
Artichokes	0.10	0.17	Trace of t
Asparagus	0.11	0.10	_
Avocados	_		Trace of t
Bananas	0.32	0.37	_
Beans, Lima	0.65	0.17	_
String, Green	0.03	0.13	_
Beets	0.11		and a second
Blackberries	trace	0.16	Trace of ox succinic, 0 cent isoo
Blueberries	1.56	0.10	Trace of o
Broccoli	0.21	0.12	_
Cabbage	0.14	0.10	_
Cantaloupe			
Carrots	0.09	0.24	
Cauliflower	0.21	0.39	
Celery	0.01	0.17	
Cherries		0.56-1.99	
Montmorency, Canned		1.45	
Corn, Sweet	_	·	
Cranberries	1.10 '	0.26	Benzoic, 0.0 cent; quinic cent (Isham
Cucumbers	0.01	0.24	_

TABLE 7.10. (Continued)

TABLE 7.10. (Continued)								
Food Items	Citric Acid (Per Cent)	Malic Acid (Per Cent)	Other Acids					
ş	2.30	0.05	Traces of oxalic					
	0.04		and succinic					
	0.34	trace						
erry	present	0.50-2.08	_					
		0.65	0.43 per cent					
			tartaric					
Concord	0.02	0.31	1.07 per cent					
			tartaric					
uit	1.33	0.08	_					
	0.35	0.05	_					
	3.84	trace						
	6.08	0.29	_					
head	0.02	0.17						
ms	_	0.14						
	0.02	0.12	-					
	0.02	0.17						
	0.98	trace	attable					
	0.13	0.35	_					
	0.37	0.37						
:d	0.05	0.69	-					
	0.24	0.12	_					
Canned	0.42	0.16	_					
esh	0.11	0.08	_					
ons, Japanese		0.09	garganite					
2	0.84	0.12						
Infornia	0.03	0.92						
η		2.48	_					
, Idaho	0.51	_	quadrates					
Cuban	0.07	_	_					
talian Style		1.44	and the same of th					
1	_	0.15	_					
ies, Black	1.06	_	_					
100, 270010	1.30	0.04						
	0.41	1.77	0.12 per cent oxalic					
	0.08	0.09						
	0.04	0.32						
10,	0.91	0.10	-					
16.	1.08	0.16						
	0.30	0.20						
	0.47	0.05						
₩hite	0,47	0.23						
inte		0.20						
	0.05							
'heat Flour rries, Canned	0.62	0.24	-					

d from Libbe 28 in Bridges, M. A. and Mattice, M. R., "Food and Beverage Analysis," Lea & 1.2 lphia, Pennsylvania, 1942.

tract of strawberry juice; butyric acid in extract of carrot seed oil; and propionic acids in molasses; and octanoic, valeric, and n-nonoic in cocoa.

Volatile Sulfur Compounds

Some vegetables have volatile sulfur compounds as an important p the flavorful materials present; others produce volatile organic sulfur pounds when the cells of the vegetable are crushed, enzymes are released hydrolysis occurs. In still other vegetables volatile sulfur compoure formed during cooking when acids are released from the cells an action with larger molecules takes place. Our knowledge of these pounds is fragmentary because few vegetables have been studied in respect and even these few not intensively.

Garlic, onions, and related species owe their peculiar penetrating and flavor to sulfur compounds. In 1892, Semmler reported that garl contains 60 per cent of diallyl disulfide, $C_6H_{10}S_2$; 6 per cent allyl p disulfide, $CH_2=CHCH_2SSCH_2CH_2CH_3$; and 20 per cent of dially sulfide, $C_3H_5SSSC_3H_5$. In more recent years investigation of garlic indithat "garlic oil" is not present in the whole garlic, but is formed on c ing. Crushing inaugurates an enzymatic reaction that releases the follows:

Stoll and Seebeck⁷⁷ have shown that the parent substance in gardalline, which forms allyl thiosulfinate on crushing. Onion has been shown to contain allylisothiocyanate, CH₂=CHCH₂CNS, as well as allyl predisulfide.

The cruciferous plants are a large family (Brassica) noted for peppery flavor and their content of organic sulfur compounds. The clude the cresses, radishes, mustards and cole vegetables as well as a weeds and garden flowers. Synge and Wood⁷⁹ have identified S-me cystein-S-oxide in cabbage and have detected it in turnip, caulific

CH₃ SO CH₂ CHNH₂ COOH

S-methyl-cystein-S-oxide

sheperds' purse, and white mustard but not in water cress, radish, or other members of the family which have been tested.

te isothiocyanates are a group of organic compounds, commonly mustard oils." The name arises from the occurrence of allyl isogranate in mustard oil. It has also been detected in many other memorithis plant family.

CH₂=CHCH₂NCS Allyl Isothiocyanate "Mustard Oil"

n distillation of radishes produces an isothiocyanate which is either putylthio)butylisothiocyanate, $C_4H_9SC_4H_8NCS$, or 4-(n-butylthio)-nylisothiocyanate, $CH_4H_9SCH_9CH=CHCH_8NCS$. β -Phenylethylisovanate has been isolated from ground water cress.



β-Phenylethylisothiocyanate

opson and Halliday⁷¹ measured the amount of hydrogen sulfide adduring the cooking of cabbage and cauliflower by condensing the ate and precipitating the sulfide ion as cadmium sulfide. The organic compounds in the filtrate were estimated by oxidizing to sulfate tremine water and precipitating as barium sulfate. They obtained denations of the rate of formation of hydrogen sulfide and organic compounds but did not identify the organic sulfides. They found that nount of hydrogen sulfide increases between 5 and 20 min in boiling and the organic sulfur between 7 and 30 min. Cauliflower gives off hydrogen sulfide and volatile organic sulfur during the same cookery ts. Since the most acceptable products are formed when cooking time volong enough for the vegetable to become soft, they believe protooking produces disagreeable flavors and odors through the formation of hydrogen sulfide and organic sulfur compounds.

evolution of hydrogen sulfide during the cooking of corn can be istrated. Since corn is not cooked for great lengths of time, hydrogen formation is no problem; but during canning, where it is important righer temperatures and longer periods of processing, the volatile formation may account in part for the "canned corn" flavor. So far botts on sulfide formation during "flash" canning of corn are

ole.

In one study volatile sulfur compounds were measured in a number vegetables. During processing a small amount splits off from peas, as gus, and kohlrabi but more from spinach and beans. The products of o kettle canning contain less volatile sulfur compounds than the fresh value table since the sulfides are rapidly vaporized from the kettle. During cooking of carrots, chanterelles (a kind of mushroom sometimes controlled tops"), and celery enough was volatilized so that none was determined to the cooked food.

The formation of volatile sulfur compounds is probably responsible the development of the cooked flavor of many vegetables and the cooked flavor of some on prolonged boiling. Some vegetables such onions become increasingly mild with cooking—probably the result of solution and vaporization of the sulfur compounds, so prominent a partheir flavor. Others such as cabbage become increasingly strong flavor which indicates that more and more volatile sulfur compounds are thesized as cooking progresses. Mellowing or ripening of flavor oc when stews and soups containing vegetables such as onions, garlic, or are allowed to stand; such mellowing is probably the result of the dis sion of these substances through the mix.

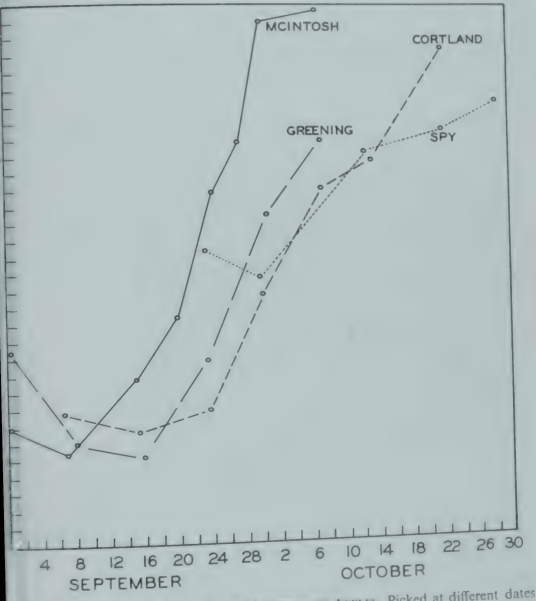
The origin of these volatile sulfur compounds is believed by some vestigators to be in glycosides hydrolyzed by enzymes, or from the vated temperatures and the release of acids during processing. Little lation work has been done, and we are still ignorant of the precursor most of these compounds. Much study is needed for a complete un standing of the role of sulfur compounds in the preparation of fruits vegetables.

POST HARVEST CHANGES IN FRUIT

It has been recognized for many years that fruit continues to under chemical changes after harvest until finally spoilage occurs as it is tacked by fungi, yeasts, or bacteria. Since the eating quality changes of these reactions and the monetary value of the crop depends on it, a fruits have been studied extensively. Both apples and bananas have the subject of many investigations, and a considerable body of knowled has been built up through the years. Pears and cherries have been studies frequently, while some fruits have scarcely been investigated. Applears, citrus fruits, and bananas are stored for variable lengths of time fore they are consumed; and the changes which occur in them are signant economically. Cherries, plums, and berries of all kinds are in perishable and reach market soon after picking. If these fruits are kept any length of time, they must be either canned or frozen.

changes of fruit after harvest are numerous and include changes in piration. (2) water content, (3) carbohydrate, and (4) organic acids

At the time of harvest the respiration rate of apples and pears has a low level. However, soon after picking, the uptake of oxygen and oduction of carbon dioxide begin to speed up until finally a climax thed, called the *climacteric*. It is followed by a steady decrease in atory rate, often called *senescence*. Pearson and Robertson have the respiration of Granny Smith apples in Australia and have that apples left on the trees for longer than 250 days past petal fall



RESPIRATION RATE OF FOUR VARIETIES OF APPLES. Picked at different dates at 74° F (23° C). Each point on each curve is the respiration rate made 24 hours ch harvest date. Courtesy of R. M. Smock.

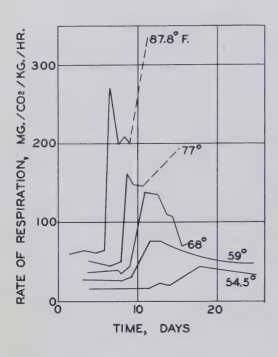


FIGURE 7.8. RATE OF PRODUCTION OBON DIOXIDE BY BANANAS (INITIALLY LAT DIFFERENT TEMPERATURES. The cur 77° F and 87.8° F are terminated by growth indicated by the broken lines. duced from Von Loesecke, H., "Banana terscience Publishers, Inc., New York from data of Gane, R., New Phytologist, 1 (1936).

(normal harvest is at 170 days) show a climacteric similar to picked Bananas are normally picked green, and they show a climacteric as ripen. Tomatoes and possibly other fruit also show this surprising chin respiration.

- (2) When fruit is picked and severed from the plant, water no leftlows into the fruit although the loss of water continues. Usually water not be taken in through the skin. In apple storage one problem is to vent water loss so that the fruit does not wither and decrease in value dry atmospheres, and particularly at high temperatures, water loss is reapples rapidly cooled after delivery to the storage barn have a smaller water loss than those cooled slowly. During the ripening periodananas, the pulp increases in water content and the peel decreases. Voloss is checked in bananas (and probably other fruit) by the waxy lay the skin. Golden Delicious apples, which are very susceptible to water have a thin skin and, what is more important, this skin is covered with and fissures through which water evaporates rapidly.
- (3) Many changes occur in the carbohydrate fraction of fruit dripening, during the climacteric, and during senescence. See Figure Green fruit usually contains an abundance of starch, but is short of soluble sugars that give ripe fruit its sweetness. On ripening, how starches decrease and sugars increase in concentration. Since these changes of the been observed, it has been assumed that the sugars are duced at the expense of the starch. However Hulme²⁶ has found the Bramley's Seedling and Early Victoria apples the loss of starch does parallel the increase in sucrose or reducing sugars. Also important in

st is the work of Barnell³ on ripening bananas: he records a steady starch: whereas his data for sugars show an increase in some cases decrease in others. See Table 7.11. These data indicate that the relatetween starch and sugars in fruit is complex. In apples and pears see, glucose, and fructose are the important sugars, with fructose the abundant. See Table 7.12. The source of these sugars and their role in polism during the climacteric have been the subject of some speculabut as yet the course of the reactions has not been established.

pectic substances are present in the cell walls, they have been the substances in the substances after harvest. Apples in storage slowly soften, he rate depends on the temperature of the storage barn as well as the sy of the apple. The graph (Figure 7.10) shows the course of softening he close correlation with protopectin and soluble pectin content. Spectin falls and soluble pectin rises until late in the storage period a reversal of this trend occurs. A decrease in chain length as well as



RE 7.9. DISAPPEAR-DE STARCH ON RIP-OF APPLES. Photoaken on August 27, I apples stained with From left: green an, ripening McInipe Wealthy. Cour-Kalamazoo Gazette.

TABLE 7.11. CARBOHYDRATE COMPOSITION OF BANANAS DURING "EATING RIPE" STAGE EXPRESSED AS PERCENTAGE OF FRESH PULP"

Heavy 3/4 Full. Ripened at Tropical Temperature							
Days from Harvest	Starch	Total Sugars	Sucrose	Glycosidio			
7	1.56	13.450	3.660	0.3			
9	0.654	12.770	1.700	0.2			
11	0.357	12.600	0.990	0.20			
	Heavy 3	/4 Full. 14 Days at 53°F, R	ipened at 68°F				
Days from Harvest	Starch	Total Sugars	Sucrose	Glycosidic			
19	7.52	6.586	1.888	0.24			
21	3.67	10.210	2.241	0.63			
24	1.62 ^b	11.245 ^b	2.356b	0.63			
	Standard	3/4 Full. 14 Days at 53°F, I	Ripened at 68°F				
Days from Harvest	Starch	Total Sugars	Sucrose	Glycosidic			
20	5.25	11.610	3.755	0.30			
22	2.89	12.720	3.890	0.20			
24	1.61 ^b	12.895 ^b	2.220b	0.2			

^a From Barnell, H. R. "Studies in Tropical Fruit XIII Carbohydrate Metabolism of the Banar During Storage at 53°F and Ripening at 68°F," *Ann. Botany*, 5, 608 (1941).

^bValues thus marked are for fruit which would be judged by the eye to be slightly overripe.

loss of methyl groups probably occurs during the softening period and counts for the rise in soluble pectin.

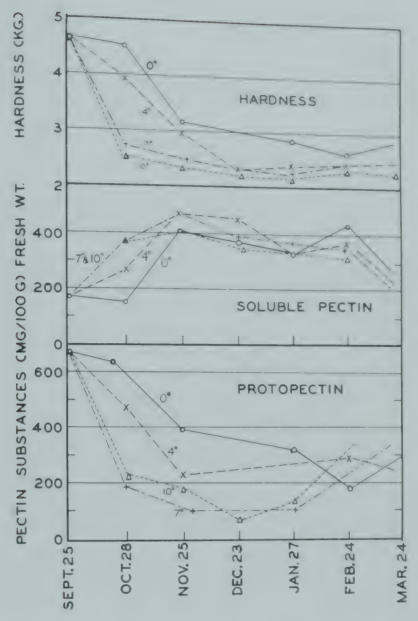
Pears are picked in the hard stage and held at low temperatures uniquired for ripening. On return to room temperature they rapidly riper soften. If pears are held too long at low temperatures, they de "sleepiness" and instead of ripening satisfactorily they turn brown as soften. In pears as in apples there is a change in the pectic substances ing softening with a rapid drop in the amount of protopectin and a r soluble pectin. See Figure 7.11. During cold storage the amount of pectin increases. Bartlett pears that develop "sleepiness" do not show rise in soluble pectins on exposure to ripening temperatures during Jar or February. It is believed that during cold storage inactivation of personness.

TABLE 7.12. SUGARS IN APPLES AND PEARS*

Apple Juice	Pear Juice (26 Varieties)	
6.6- 56.6 g/L	1- 24 g/L	
12.3- 58.0	5- 35	
69.2–113.8	65–112	
	6.6- 56.6 g/L 12.3- 58.0	

^{*}Data from Tavernier and Jacquin, reported by A. C. Hulme, "Some Aspects of the Biochemi Apple and Pear Fruits," *Advances in Food Research* **8**, 297–413 (1958).

RE 7.10. CHANGES RDNESS, SOLUBLE AND PROTOPECTIN. Two in fruit for apples during storvarious temperadapted by A. C. rom Ulrich.



nes slowly occurs and normal hydrolysis of protopectin does not

nanas undergo loss of protopectin and increase in the smaller molewhich make up soluble pectins during ripening. Von Loesecke⁸⁴ has nted data (Table 7.13) for various varieties of bananas.

s of protopectin and rise of soluble pectins have been found in soluble. It is rlums, and tomatoes as they ripen. Raspberries do not show a fation between the pectic substance fractions and the development of mess in the fruit as it passes its peak of ripeness.

In the fruit as it passes to peak in the

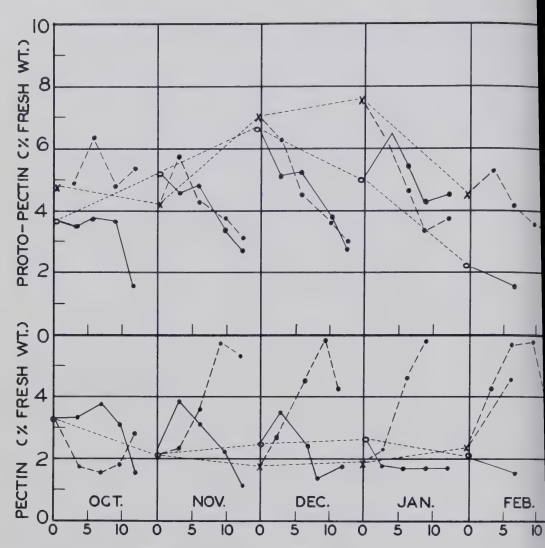


FIGURE 7.11. PECTIC CHANGES IN BARTLETT (0) AND ANJOU (x) PEARS. Stored -1.11°C to -.56°C and removed at monthly intervals for ripening at 20°C to 21.1 Dotted lines show changes during cold storage while interrupted lines (Anjou) and continulines (Bartlett) show changes occurring in the ripening room. Adapted from Date Hansen, *Proc. Indian Acad. Sci.*, **B39**, 17 (1954) by A. C. Hulme, *loc. cit*.

xylans and arabans and a decrease in cellulose. Pears contain lignins in stone cells and are unlike other fruits in this respect. Hulme does not consider a few studies on apples definitive for hemicellulose. The alcohol soluble fraction of apples is partially soluble in dilute acid and containing with pectic substance a so-called "hemicellulose fraction." This fittion slowly decreases during storage.

(4) Organic acids decrease in apple pulp and in pears during stora Both of these fruits contain a large number of organic acids in low c centration, a small amount of citric acid, and larger amounts of malic ac With the development of the techniques of chromatography, it is now p

ABLE 7.13. CHANGES IN PECTIN AND PROTOPECTIN IN PULP OF BANANAS*

(Per Cent of Fresh Pulp)

		0	3 Da	ys in Ripening 5	Room 7	9	11
Michel	Pectin	_	0.27	0.36	0.34	0.37	0.40
Michel	Protopectin	0.53	0.56	0.31	0.34	0.21	0.22
	Pectin	0.21	0.48	0.52	0.57	0.58	0.68
Finger	Protopectin	0.50	0.76	0.27	0.29	0.31	0.35

Subjished data from Manion, J. T., United Fruit Co., Research Department, 1933 in von Loesecke Bananas, Chemistry, Physiology, Technology, 2nd ed. Interscience Publ., Inc., New York, N.Y.

to separate and identify compounds present in low concentration were extremely difficult to find by other classical methods. The techhas not been widely utilized to follow the changes in concentration rious acids, but some information has been obtained. Quinic acid and mic acid are evidently much more widely distributed in plant tissues has previously been recognized. Hulme²⁶ reports that in storage at Bramley's Seedling apples show changes in total acids, with a differattern for different acids in pulp or peel. (Table 7.14) The changes in ic acids during storage should be of considerable interest, since organic acids are related to metabolic processes. The Krebs cycle for retabolism of two carbon fragments has not been demonstrated comy in apples and pears as yet, but the possibility of its importance is

TABLE 7.14. CHANGES IN ORGANIC ACIDS IN BRAMLEY'S SEEDLING APPLES STORED AT 75°C*

	SEEDLING AFFEES 5	TORED AT 70 C	
	15 Days mg/100 g	40 Days mg/100 g	100 Days mg/100 g
ic	6-8.5		10
	fresh tissue	80	50
nic	45	-	1-2
cimic	_		
			1-2
6	1–2	400	
stic	400	400	8
timic	5	10 at 25 days	25
ımalic		(A b. and Pear Fruits	Idvances in From

in Hu me. A. C., "Some Aspects of the Biochemistry of Apple and Pear Fruits. Advances in Food th. 8, 329 (1958).

great since it is used by many forms of life as a metabolic pathway. function of acids such as quinic and shikimic is as yet unknown, but surely have some role in metabolism.

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1ilk and Milk Products

AND MILK PRODUCTS have formed an important part of the diet of tern man since the dim reaches of history. It is not known when aniwere first domesticated but soon after man became a farmer, he began ep and raise animals. All mammals produce milk after the birth of the ig and man has used the milk of many animals for his own food. The is, of course, the most important of all these animals as a supplier ood for man, but yak, reindeer, buffalo, or goat milk is more imant in some parts of the world.

r many generations numerous families kept their own cow and the was supplied fresh from the udder, morning and night. However, the rise of cities and the growth of specialties, the dairy industry oped. Both governments and dairies attempted to develop checks on urity of milk, both from the chemical and bacteriological stands. The United States has extensive federal regulations on the milk and products that pass in interstate commerce, while each state has many for the regulation of their production and sale within its boundaries.

e interest in obtaining and producing high grade milk and milk prodnes resulted in a large number of studies of the composition of milk. tudies for the most part have been slanted to answering the problems ntrol, but nevertheless much general information about milk and milk icts has accumulated.

the United States when the term "milk" is used, it always refers to the of cows. If the milk of another species is intended, the name of that is precedes "milk"; i.e., "human milk" (sometimes "woman's milk"). milk," etc.

od and Drug Administration of the U.S. Department of Agricul-Irlined milk in August, 1926. "Milk is the whole fresh, clean, lacteal ion obtained by the complete milking of one or more healthy cows. rl fed and kept, excluding that obtained 15 days before and 5 days

after calving, or such longer period as may be necessary to re milk practically colostrum free."

FORMATION OF MILK

The production of milk by the mammary gland of any female depends on three phases; (a) the development of the mammary glasecretion of milk, and (c) emptying of the gland. After conception implantation of the embryo into the uterine wall, a series of chang in the mammary glands. These are under the influence of he elaborated by the ovary and the corpus luteum. The mammary glasgin to proliferate; the epithelium of the lobes increases rapidly secretory cells develop. Toward the end of pregnancy these gland to form a secretion. When parturition occurs, the influence of the phormones is removed from the pituitary, permitting it to form a he prolactin, which causes secretion in the prepared mammary gland.

The first secretion of the gland differs from milk and is called conthe composition of which is compared below with that of milk. It atively thick fluid during the first days postpartum, gradually charappearance and composition to those of milk. It differs from milk that its protein content as well as some of the vitamins and amino Rather recent work with cow's colostrum gives the following compared with normal milk. [Quoted from Sutton's report in Macy Kelly, H. J., and Sloan, R. E., "Compositions of Milk," Natl. Ack Natl. Research Council, Publ. (1953)].

Per Cent of Total Nitrogen

	Casein	Albumin	Globulin	No
Colostrum	31	6	55	
Mature Milk	71	8	18	

In a summary of the work to 1953 on the vitamin composition two fluids, Sutton points out that vitamin A activity is much his colostrum than in milk. The concentration is 10 to 15 times higher the first few days of secretion than the average for milk. Likewis flavin is present at a 3 to 3.5 fold concentration. Colostrum is his thiamine and biotin, about the same in niacin, and lower in pantacid. It also contains more of the amino acid, tryptophan, than milk.

In women it is common to consider the period of colostrum for the first five days postpartum; of transitional milk formation, days and mature milk formation after the tenth day. Actually, evidence in that the period of colostrum formation differs with individuals and I to 5 days. Cows and goats have a shorter period of colostrum for-,111.

the secretion becomes characteristic of that of milk, its comon remains relatively constant. It continues to flow for some months est species if the gland is emptied regularly. If the milk is not reed, it disappears, and the breast tissue undergoes involution and reto the pre-pregnancy state.

APOSITION OF MILK

Jk is a complex mixture of lipids, carbohydrates, proteins, and many organic compounds and inorganic salts dissolved or dispersed in . Macy, Kelly, and Sloan list over 100 compounds in it. Some of compounds such as the carbohydrate, lactose, and most of the salts sitamins are soluble in water. Others such as the lipids, proteins, and cium phosphate are dispersed through the water in the colloidal or colloidal state.

e lipid is composed primarily of fat although there are also small unts of phospholipids, sterols, the fat soluble vitamins A and D, caroand xanthophyll. The proteins (see p. 140) are numerous and are tionally classed in the following fractions which do not consist of pure uns: (1) casein precipitated by rennin or acid, (2) lactalbumin, and (3) globulin precipitated by heating. The carbohydrate of milk, whether cow or other species, is lactose and it is the only carbohydrate preshe ash of milk contains all of the minerals which occur there, but not sarily in the same form. The salts are not only those of inorganic acids frome organic ones such as citric acid as well. Part of the phosphate 's as phosphoprotein and phospholipid, part combined with calcium. he ash of milk which has been most extensively studied rather than unerals in the milk itself. Calcium phosphate is known to be colloiddispersed in the water (it has a very low solubility in water), and it own that milk contains potassium, sodium, magnesium, and chloride Il as small amounts of copper, iron, zinc, manganese, aluminum, and e. Sulfur is present in some of the amino acids of the proteins.

avitamins are plentiful in milk. Only ascorbic acid is present in limmounts, and it may even be absent from pasteurized milk. But the sat r and fat soluble vitamins are there in goodly quantity. The B a vitamins are dissolved in the water and vitamins A and D in the inter milk is not a rich source of vitamin D unless the diet of the

k contains a number of enzymes, some of them apparently secreted nilk and some of them formed by the microorganisms which inhabit

TABLE 8.1. COMPOSITION OF MILK

	Range (%)	Average (%)
Water	82.0-90.0	87.3
Fat	2.3- 7.8	3.67
Total Protein	2.0- 4.5	3.42
Lactose	3.5- 6.0	4.78
Ash	0.6- 0.9	0.73
Solids	10.0-18.0	12.69
Solids-not-fat	7.5–10.6	8.77

From Jacobs, M. B., "Milk, Cream, and Dairy Products," in Jacobs, M. B., ed., "The Chemist Technology of Food and Food Products," 2nd ed., Interscience Publishers, New York, N. Y., 1951.

the milk. Amylase, lipase, tryptase, peroxidase, catalase, and reduchave all been demonstrated. Galactase, lactase, and aldehydase have reported. Some of the enzymes are destroyed or reduced on pasteurization and tests for their presence or level may be used as a test for effect pasteurization.

The milk sold on the consumer market is pooled milk from many h and consequently is relatively constant in composition. The ranges in c position are given in Table 8.1. Variations in the composition of milk cur with breed of cow, time of year, time of day, portion of the milk time in the lacteal cycle, and the nutritional status of the cow.

Some breeds of cows give milk which is always higher in butterfat that of others. See Table 8.2. Thus Jerseys and Guernseys are noted for high percentage of fat. The protein of their milk is likewise a little high than most other breeds, and the percentage of water is consequently low

The time of year has an effect on the composition of the milk although this may be indirect, since the diet of the herd and the amount of grasture consumed varies with the time of year. Jacobs⁴ presents ta

TABLE 8.2. VARIATION IN COMPOSITION DUE TO BREED

Breed	No. of Cows	Fat %	Protein %	Lactose %	Ash %	Total solids %	Water %	So No
Jersey	29,495	5.37	3.92	4.93	0.71	14.91	85.09	9
Guernsey	32.562	4.95	3.91	4.93	0.74	14.61	85.39	9
Brown Swiss	721	4.01	3.61	5.04	0.73	13.41	86.59	9
Ayrshire	6,999	4.00	3.58	4.67	0.68	12.90	87.10	8
Shorthorn	6,155	3.94	3.32	4.99	0.70	12.81	87.19	8
Holstein	37,598	3.40	3.32	4.87	0.68	12.26	87.74	8

From Jacobs, M. B., loc. sit., p. 848.

n Gamble, J. A., Ellis, N. R. and Besley, A. K., U. S. Dept. Agr. h Bull. 671, (1939), for both Holstein and Jersey cows which show lations in the amount of fat, protein, lactose, ash, iron, copper, calcium, phosphorus. The variations are not too regular. Protein in Holstein for January through December shows the following monthly values: 1, 3.09, 3.08, 3.16, 3.14, 3.10, 3.06, 3.19, 3.18, 3.30, 3.34, 3.17. Woodis says that the poorest milk is produced in the spring and early sumwhen green pasture is bountiful and the richest in autumn and winter. Table 8.3. However, the table from Gamble, et al. that was previously stioned does not bear out this trend. The fat for Jersey milk from ruary through December showed the following monthly values: 5.1. 5.5, 5.4, 5.5, 5.4, 5.2, 5.0, 5.5, 5.2, 5.5 while for Holstein: 3.8, 3.7, 3.7. 3.2, 3.4, 3.5, 3.2, 3.5, 3.6, 3.5. September is the low month for both eds, but there is no regular pattern through the year.

he composition of milk varies with the time of day and portion of the king. The evening milk tends to contain more butterfat and slightly less er than morning milk. Likewise the milk which is first removed from the er contains a smaller amount of fat than the "strippings" that re-

ved in the final phase of milking.

he time in the lactational cycle, i.e., the time after calving also has an sence on the composition of the milk. For example Duncan, Watson, in and Ely3 found that the threonine level in milk proteins is 15 per t higher at the end of the lactational period than 60 days after birth of calf.

he diet of cows has a very marked influence on the quantity of milk duced, but only a limited effect on the composition. Diets have been ely studied, but complete knowledge of all of the factors that influence x production is still in the future. Any dietary deficiencies of protein, al energy, etc. are immediately reflected in a decrease in the total volume nilk formed. It is true that the vitamin D and carotene content of the reflect the level in the cow's diet, but for other components the quan-

TABLE 8.3. VARIATION IN MILK WITH SEASON

Time of Year	Total Solids	Fat	Solids-Not-Fat
ber-January	13.04	4.11	8.93
aary-April	12.72	3.88	8.84
August	12.66	3.89	8.77
ber-November	13.03	4.25	8.78

m Woodman, A. G.8 (Data of Richmond).

tity present in the milk is relatively constant, and it is the total secret all components which is influenced by a lowered intake.

Diet does have an effect on the constituents of the milk responsib the development of "oxidized flavor." This is a flavor which is various scribed as "cardboardy" or "oily" and it may be so marked as to make milk completely unpalatable. It appears to develop when milk is in co with oxygen and particularly rapidly if the milk contains small amour copper. Milks vary considerably in their tendency to develop this f and a fair amount of work has gone into attempts to understand and trol it. Various compounds have been implicated as the active age phospholipids, ascorbic acid, and the tocopherols—but so far it is no equivocably known. Krukovsky, et al.6 have shown that diet has a Cows fed roughage from late cut hay, timothy in full bloom, and all whether fed as silage or as field cured hay, did not produce a single ple of milk which developed an oxidized flavor. The milk was teste adding 0.1, 0.5 or 10 mg copper per l of milk and holding the sample 0°-5°C for 10 days. However 53 per cent of the samples from cows early-cut hay, either silage or dried hay, developed oxidized flavor-s of them very strong.

Milk possesses a fine delicate flavor that results from the blendin many compounds which affect taste and olfactory endings. A flavor di ent from that of normal milk is objectionable and milk with an unu flavor is not accepted by the consumer. Josephson⁵ discussed flavor milk in his American Chemical Society Borden Award address and tributes the "sunlight" flavor which is produced when milk is expose sunlight to the photolysis of the amino acid, methionine. A "cowy" fla occurs when acetone bodies appear in the milk. Cows are prone to dev ketosis, a condition in which fatty acids are not completely oxidized carbon dioxide and water and in which some acetone bodies are form The acetone bodies are acetone, β -hydroxybutyric acid, and acetoa

CH₃COCH₃

CH,CHOHCH,COOH

CH, COCH, CO Acetoacetic A

Acetone β -Hydroxybutyric Acid

acids. When they are produced in an animal, they are carried by the bl and excreted in urine and breath. In a lactating animal they also appear small amounts in the milk.

The flavor of milk changes when it is heated for any length of time as "cooked" flavor develops. This is noticeable in evaporated milk even w great precautions are used to avoid overheating. Josephson attributes "cooked" flavor to heat denaturation of lactalbumin. He attributes elized" flavor which sometimes develops to changes in lactose with rmation of furans.

cr of cows. Many dairies produce "standardized milk" which conapproximately the same amount of butterfat throughout the entire Usually the standard is that required in the particular area served adiry, and it can be met by adding cream or skimmed milk to the of this is allowable. The regulations governing the sale of milk are stringent. Even when a dairy serves a relatively small area, its prodoften subject to local laws. These laws usually have some regulations ding microorganisms and filth, as well as some regarding butterfat int. Many state laws allow standardization by the addition of either to reskimmed milk.

bacteriology of milk is very important. In the udder, milk is related of free of microorganisms, bacteria, molds and yeasts, but it is readily minated by organisms on the outside of the udder, on the hands of the ror the milking machine, and on the vessels in which it is transford and stored. Not only is milk an excellent food for the nourishment n, but it is also fine for the nourishment of microorganisms. When the introduced into milk, they flourish and multiply rapidly. A food tharge population of microorganisms is not only worthless as food ghly dangerous.

AICAL ANALYSIS OF MILK

past milk has often been bought on the basis of its butterfat confine practice of paying for the milk solely on the basis of fat, if it up to standards as far as bacterial count, is not as common as rly, but the amount present is still determined. Every little dairy in untry has the simple equipment necessary for the determination of tat. The Babcock method is used widely for this determination and gn it is not too simple to carry out with accuracy, it is used by many who have had no training in chemistry. Some states have legal rems on the equipment used in the determination, in order to insure cy in able hands.

Babcock method is one which releases the fat from the milk emuld measures the percentage directly in bottles calibrated for this. A and volume of milk, 17.6 ml is treated with 17.5 ml of concentrated chick. The acid denatures and partially hydrolyzes the protein, so no longer acts as a protective colloid. The fat rises to the top. It is kept liquid by heat from the reaction and by warming the milk-aciture, as the determination proceeds. The bottles are whirled in a cer which causes the fat to rise. Hot water is added and the process is re. The fat which accumulates in the neck of the bottle and which is percentage butterfat is "crude fat" and contains all the lipid mater the milk.

The Röse-Gottlieb (because of the umlaut, sometimes spelled method requires extraction of the milk with a mixture of ethyl eth petroleum ether, distillation of the solvent and weighing the fat r. The fat is then dissolved in petroleum ether and any insoluble in weighed. The difference gives the butterfat content of the sample. also a "crude fat" determination since it includes all compounds are soluble in petroleum ether. This comprises all lipids, the fat s vitamins, and pigments.

Milk solids include all of the compounds present in milk except and are readily determined by evaporation of a sample in a weigh bottom dish at approximately 100° C to constant weight. It is necess avoid heating the sample above this temperature since browning residue will occur. Milk solids can be determined approximately by uring the specific gravity with a Quévenne lactometer and applying the mula (%Solids = 0.25L + 1.2F + 0.14) where L = 1 lactometer reading must be rected to 60° F before the formula is applied. The total solids of magnify constant and this is a useful determination in indicating some of adulteration. Often the solid-not-fat is calculated by subtracting amount of crude fat from the total solids. Since fat shows wider flations than the other components, this value has slightly more contain total solids.

Lactose may be determined by a number of methods. Those which commonly used are not specific for lactose but since it is the only bohydrate present in milk, they serve well to give fairly accurate a producible values. Either the determination of the optical activity reducing capacity with a copper reagent are used. In the determinate the optical rotation, the protein is precipitated and the filtrate reappolarimeter. The assumption that lactose is the only remaining computate possesses optical activity is not quite valid since some of the components are optically active but are present in very low concentrations.

The copper reducing method is also applied to the protein-free fi The proteins are commonly removed by adding copper sulfate to t luted milk and precipitating most of the cupric ion with either sodi potassium hydroxide. The cupric hydroxide which forms carries down sein as well as the fat. An aliquot of the filtrate is treated with I chless Solution and the cuprous oxide formed either filtered off, dried, and ghed or determined by one of the other standard methods.

roteins of milk are ordinarily determined by the Kjeldahl method (see thod for total proteins in milk suffers from the same limitations as does use of the method for the protein content of any food. It will be noted the constant used for milk differs slightly from the general constant, defractions, not pure proteins. Casein is precipitated by warming the nitrogen by the Kjeldahl method and the percentage of nitrogen denired is multiplied by 6.38. The albumin is precipitated from the ate obtained after removal of casein, by neutralization, addition of the rect amount of acetic acid and heating on a steam bath. This precipities likewise analyzed for nitrogen by the Kjeldahl method and the centage of nitrogen multiplied by 6.38.

asteurization is an important process in insuring that some of the roorganisms are destroyed. A number of chemical tests have been ded to determine whether or not a given sample of milk has been teurized or if the process has been carried out satisfactorily. Enzymes sensitive to heat and some of them are denatured by the pasteurization. teurization is carried out by heating the milk to 142 to 150°F for 30 utes or by heating it (flash pasteurization) to 160°F and holding for 15 ands. Although this is a relatively low temperature, most of the entes of milk are destroyed. Tests for the enzymes can therefore be used as emical means of differentiating raw from pasteurized milk or of detectincompletely pasteurized milk.

he Schardinger test detects the presence of peroxidase in milk. An alolic solution of methylene blue, formaldehyde and water is added to m.lk samples. The tubes are placed in a water bath at 45°C. In less 120 minutes the raw milk will decolorize the methylene blue while the carried milk will take much longer.

he Kay and Graham test for phosphotase is more delicate and can ly detect faulty pasteurization procedures. A sample of milk is intend with phenyl phosphate in a diethyl barbiturate buffer for 18 to 24 at 34 37°C. In the presence of phosphatase the phenyl phosphate drolyzed and phenol is formed.

$$C_6H_5OPO_3H_2 + H_2O \rightarrow C_6H_5OH + H_3PO_4$$

e milk has been pasteurized sufficiently, most of the phosphotase will

have been destroyed and little hydrolysis will occur. The phenol is termined colorimetrically and any values over 0.047 mg phenol/0.5 milk indicate progressively inadequate pasteurization.

CHECKS FOR PURITY

The constant vigilance of the governmental agencies as well as the velopment of dairying into a big business anxious to maintain quality, I resulted in pure milk for the consumer in the United States. Milk sol unadulterated chemically and fairly pure bacteriologically. (Absolute st ity except in canned products is impossible.) In generations past and some parts of the world today this is not true. In some instance there been an addition of water and skimmed milk to increase the volume w milk has been sold on a fluid basis, of thickeners such as gelatin calcium sucrate to make cream appear richer, and of preservatives coloring matter. During the past 50 years a large literature has develo on these adulterants and methods of detecting them.

Since the composition of milk is variable, it is difficult to detect waing, skimming or the addition of skimmed milk. Watering is illegal in states of the Union, but standardization is permissible in many. The action of water decreases the percentage of all components, but if the amo added is small, it may simply reduce them to the lower limits of the components. Fat shows the greatest variation in milk and consequently determination of the solids-not-fat is frequently used as an index of waing. In Table 8.1 the range for solids-not-fat is given as 7.5–10.6 per composition of the value for a sample of milk is close to the bottom of this range would surely be suspicious.

Milk serum or whey, that portion of the milk from which both fat a casein have been removed, is quite constant in composition and analysis it is often used in attempting to detect watering. The refractive index, specific gravity, or the total solids are determined. If the milk has b watered, the solids are present in lower concentration and the refracting and specific gravity will consequently be closer to those of water the for pure milk.

One of the most reliable physical constants of milk is its freezing po The presence of dissolved substances, the salts, the lactose, and ot molecules, depresses the freezing point of milk below zero while colloi substances have a slight effect on the freezing point. Pure milk freezes tween -0.530 and -0.566° C with an average value of -0.545° C. Milk wh has been watered will have a freezing point closer to zero. Here, too, range in value does not allow a sharp differentiation. A small amount r might be added to milk which contained an unusually large amount sluble substances without raising the freezing point above the normal

imming of milk or the addition of skimmed milk to ordinary milk does hange the amount of any of the components except fat. It can be deat, by calculating the ratio of the fat to protein, lactose, solids, or solidsat. But here again, since the ratio of these constituents is not constant flerent samples of milk, this is difficult to prove. If the fat falls below tandard set for that area of the country, but the protein, or solids do then skimming has doubtless been used.

eservatives in milk are prohibited by law. The milk delivered must be in fresh condition by the use of sanitary procedures, equipment, packand adequate refrigeration. In the early days, preservatives were compused to extend the time during which the milk was saleable. Formade in very small quantities has the ability to preserve the milk; boric or borax, salicylic acid, benzoic acid, hydrogen peroxide, and fluorides all been used. Tests for all of these preservatives and methods for deag them even when present in small amounts, have been widely used in ast.

ickening agents and coloring matter have more frequently been added cam than to milk. Cream is expected to be slightly viscous and vellowfit is chilled and has a relatively high fat content. The addition of in calcium sucrate, or agar-agar has sometimes been used to create lusion of a higher fat content than is actually present. The use of thickeners, as well as coloring matter, is prohibited in milk and in. In chocolate drinks, egg nog and prepared milk drinks they are permissible and are sometimes used.

CIAL MILKS

ost American dairies sell more than one kind of milk today and some quite a list of products. The names of some of the more common and a very brief description of their special composition or characterwill be given.

rt fied milk is milk which reaches the bacteriological standards of the rican Association of Medical Milk Commissions. It is produced under conditions from cows known to be free of tuberculosis and brucel-ind has a low count of bacteria. Most certified milk was formerly sold that much of it is now pasteurized. Sometimes it is designated as ied-raw and certified-pasteurized.

intogenized milk is milk in which the size of the cream globules has

been reduced sufficiently so that the cream does not separate out hours. The milk is forced through very small orifices or between which mechanically reduce the size of the globules.

Vitamin D milk is widely sold and is the most common fortifie product. Milk has a relatively low content of vitamin D, particularly winter. Summer milk may have approximately 30 International u each quart but in the winter, the content falls very much below this Since the need for vitamins D during infancy and early childhood is vitamin D milk was introduced in 1932 and is widely sold now United States. The vitamin D content of milk can be improved by radiation with ultraviolet light, (2) addition of a vitamin D concentra (3) feeding a diet high in vitamin D to the cow. When ultraviolet comes in contact with some of the sterols (but not cholesterol), a che reaction occurs with a change in the ring structure of the sterol. Th product possesses vitamin D activity in young animals. The sterol wh present in milk is 7-dehydrocholesterol, the same sterol present in h skin, and it is changed by ultraviolet light to the vitamin which is nated Vitamin D₃. Irradiated milk usually contains 135 to 200 In tional or U.S.P. units per quart. When a concentrate is added to milk usually added so that the level of vitamin D is 400 International or U units per quart. Milk produced from cows which are fed a special sour vitamin D, usually irradiated yeast, produces milk which is often of "metabolized milk" since the added vitamin D passes into the milk d the metabolic processes of the cow. Where this procedure is used, the c given sufficient vitamin D so that the level in the milk produced is International units per quart.

Today almost all of the Vitamin D milk produced in the United Statortified by the addition of a concentrate. If milk is irradiated suffictoraise the vitamin D content to 400 U.S.P. units, off flavors develop milk with lower levels has not been able to compete with fortified Also since the idea of adding a concentrate has been accepted by the Alican Medical Association, this far easier method which does not respecial equipment or special care to avoid contamination has been wadopted. Metabolized milk has likewise largely disappeared from market, since the amount of vitamin D consumed by the cow is far grathan the amount which appears in the milk.

Fortification of milk with other vitamins is now practiced in many at The milk is sold under various names and is usually several cents a composed more expensive than ordinary milk. The vitamins used are often the B plex, or some of the B complex, as well as ascorbic acid and vitaming Some nutritionists are enthusiastic about this type of fortification, many are very much opposed. Their opposition rests on the widespears of the solution of th

pution of these vitamins and the ease with which they are provided in 11-rounded diet.

fled milk is skimmed milk to which some fat such as coconut oil has idded before evaporation and canning. In some states the sale of filled is prohibited.

constituted milk is milk produced by the addition of water to milk ter, skim milk powder, evaporated or frozen milk or some combinaof them. During World War II and since, reconstituted milk has been in many parts of the world where the supply of fresh milk is either scient or unsafe. It is made up so that the fat and the solids-not-fat the concentrations in fresh milk. In some states its sale is illegal.

any fermented milks are available in the United States, made by the ion of cultures of microorganisms to the milk. These organisms grow eed on milk components and form acid. Buttermilk is produced either whole milk or from skimmed through the action of the bacterium tococcus lactis. The raw milk is heated to kill most of the micronisms present, is cooled and inoculated with a culture. Incubation for ours allows the organism to grow and form acid. Buttermilk is some-, stabilized with a small amount of gelatin so that separation of the n does not occur.

idophilus milk is prepared in a similar manner by using Lactobacillus philus. The milk is sometimes more thoroughly sterilized than with ary buttermilk. Twenty or thirty years ago, there was quite a fad for nilk since it was believed that the flora of the intestinal tract has a und influence on the health of the individual and that the establishof L. acidophilus would lead to improved health. It has been shown he identity of the bacteria present in the large intestine can be changed ntinued ingestion of a large number of organisms, such as L. acidoph-But it has not been proved that the health of the individual depends

re recently yogurt has been one of the foods extolled by the faddists. old in many areas of the United States. It is an excellent food but not or to many other milk products. It is also called "bulgarlac" or oon" and is likewise a fermented milk. In this case the organism to the partially sterilized milk is Lactobacillus bulgaricus. Both chaus and bulgarious milk usually have a smaller amount of acid than milk and a creamier texture.

ER

ter is a milk product composed principally of fat. It must contain at 30 per cent crude fat and is sold with or without the addition of salt and with or without the addition of coloring matter. The 20 per cent fat is composed of approximately 16 per cent water, 2.5 per cent salt, 1.5 per cent milk solids. While neutral fat is the principal ingredient of "fat," phospholipids, sterols, pigments, and fat soluble vitamins are present. A small amount of lactic acid is also present in butter and tributes to its flavor. The amount is controlled by washing the butter, high an acid content is undesirable but if the amount is below 0.02 per the butter is more liable to putrefaction. The manufacture of butter is cussed briefly on p. 51.

Butter must contain at least 80 per cent crude fat and be free of filth, scored on a grading system which allows the following points: flavor body, 25; color, 15; salt, 10; and package, 5 to give a total of 100 pos points. Butter which scores 93 or 94 must be very mild, sweet and free flavor. Only the most expensive butter on the consumer market scores high; most is 92 score and the cheaper grades are below this. Any be scoring below 75 is called grease and is considered unfit for food. On letter scale AA is U. S. 93 score; A, 92; B, 90; C, 89; and below 89, Co cooking grade.

The addition of salt to butter inhibits the growth of some microor, isms and kills others, so that keeping quality of salted butter is better to for unsalted. However the relatively high moisture content of butter the presence in the water, not only of salts and vitamins but also of tein, make it a fine nutrient medium for some microorganisms. Butter he relatively short life unless it is stored at low temperatures. Frozen but can be kept for some months without serious deterioration in flavor. But grease which is prepared by melting butter and separating the fat from whey, has a much longer storage life, since the moisture content is very large.

CHEESE

For hundreds of years cheese has been made by man from the curd cipitated from milk by acid or rennin. The curd is given various to ments—heating, pressing, the addition of seasonings—and many che are allowed to ripen before they are used. The ripening process is on which microorganisms—sometimes bacteria, sometimes molds—grow use the curd products for food. These microorganisms form gases, and other compounds that produce flavors characteristic of the partic cheese. They also modify the protein so that the texture is altered and comes characteristic of the cheese. In centuries past when nothing known of microbiology, techniques developed so that cheeses were duced in various regions of the world that, from year to year and f kitchen to kitchen, were approximately similar. These techniques developed

ough happenstance and through trial and error. The cheeses of one type not always exactly identical but they are so similar that expectations texture and flavor of say, Gorgonzola, or Cheddar are possible. Not also the strain of microorganism responsible for the flavor developed. It is the strain of microorganism responsible for the flavor developed. It is the type of milk and its cream content. Some cheeses are made in skimmed, others from whole milk and some have added cream. Cow's less the chief milk used but sheep and goat milk are used for some es. In order to achieve fairly reproducible results, the techniques all not the line must be standardized. The curd must be treated the same ty, the ripening must be at the same temperature and for the same gth of time, etc. Considerable skill and attention to details are required produce repeatedly cheeses with similar textures and flavors.

mposition

since the curd used for the preparation of all cheeses is a casein preutate, they are rich in protein which varies from approximately 20 to 30 cent. The exceptions are the cheeses which are very moist and which requently have a slightly lower percentage of protein. Other comients vary considerably in their levels depending on the ingredients used. Varies from 1 per cent for skimmed-milk cottage cheese to 38.3 for rerican red, 39.9 for French demi-sel Cream, and 51.3 for Ricotta tta. Other components do not show this very wide range of values, but re are nevertheless differences and there are also some similarities. Most eses except cottage and English cream cheese are excellent sources of rium.

ripened Cheese

inripened cheeses are cream and cottage cheeses. Cream cheeses are preed by coagulating the casein in milk with rennet after it has soured htly. An inoculation of a culture of bacteria is added to the milk and a rt incubation period allowed before the rennet is added. The precipitate a tine creamy texture, hence the name. It may or may not be made from tm. Cottage cheese (also called, Cup, Dutch, Clabber, Pot and Smeers) is prepared from either whole or skimmed milk in which the incubathas been carried on long enough to develop more acid (0.65 to 0.7 per pressed as lactic acid). Sometimes rennet is used to complete the preation and sometimes the acid alone is the agent. The curd formed is ted to a temperature of about 120° F, the whey drained and, after ling and salting, the curd is "creamed" by beating. Sometimes cream is ed at this point.

ennet is an enzyme preparation widely used for the precipitation of

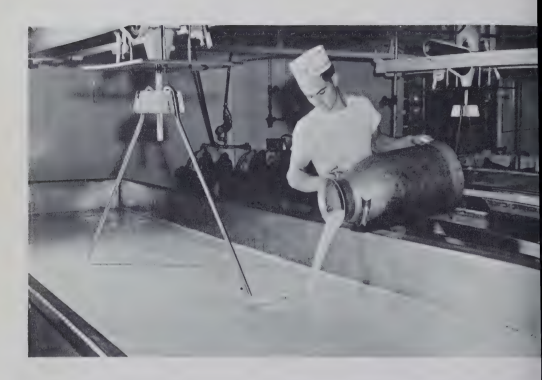


FIGURE 8.1. STARTER CULTURE. One of the agents that causes the milk to coag added to the vat as paddles rotate to stir it thoroughly into the milk. Stainless steel this type hold 18,000 lbs of milk—enough to make 1800 lbs of cheese.

Courtesy of Kraft



FIGURE 8.2. AFTER A CURD HAS FORMED. It is cut with stainless steel wire knive small cubes, to help expel some of the whey. Following this, automatic paddles will scontents of the vat while it is heated, to shrink and firm the curd and expel more who

Courtesy of Kraft



URE 8.3. THE MATTED OR CHEDDARED CURD. It is cut with stainless steel knives abs. These will be piled one on top of the other two deep, turned, and piled three using the curd's own weight to matt it out into thinner slabs.

Courtesy of Kraft Foods.



RY 84 THE SALTED CURD. It is being packed into stainless steel cheese hoops, ith heavy parchment that will form the wrapping for the finished block of cheese. ent years, round hoops were used and each was lined with cheesecloth. From here eses go to the press, where they are held overnight.

Courtesy of Kraft Foods

casein. It is a mixture of the enzymes pepsin and rennin and is used a extract or occasionally as a powder. It is prepared by extracting the li of the fourth stomach of calves with sodium chloride. In the presence calcium ions, rennin forms a thick gelatinous curd of casein. See p. 141 a short discussion of the reaction. The casein curd formed either through the action of rennet or acid is the basis for all cheeses.

Ripened Cheese

A good example of a ripened or cured cheese is Cheddar. This ye cheese was first made in Cheddar, England and now accounts for n than half of the cheese manufactured in the United States. It is sold up many names—Cheddar, American, "store cheese," the name of the comunity where it is made, or even "rat trap" cheese. Although the final concernities of other ripened cheese may be quite different from Cheddar, methods of preparation are very similar. Differences in hardness and a sistency of the cheese are the result of slight difference in the tempera at which the curd is formed, the amount of heating it receives and amount of evaporation allowed during curing. Differences in flavor also texture and consistency result from the enzymes and microorgani introduced into the cheese and developed during ripening. Some of the are present in the milk used, others are introduced in the culture, and more come from the air of the cheese factory.

Cheddar cheeses are produced by adding a "starter" to milk in orde produce acid. The culture commonly used for butter manufacture cont. ing Streptococcus lactis, S. citrovorus, and S. paracitrovorus is often u When the acidity calculated as lactic acid is approximately 0.2 per (lactic acid is not the only acid present but is the most abundant), rer and sometimes color is added and the curd forms. Great skill is required handling the curd so that the maximum yield is obtained and so that characteristics necessary to a desirable cheese are developed. The curheld until the proper degree of firmness results, a matter of 6 to 8 ho and during this period a rapid increase in acid occurs. The curd is cur form small cubes and then gently stirred to separate the whey. The mixt is then heated to 100° F. The curd is filtered and the final amount of w is removed by cheddaring or matting. This consists of rotating and turn the slabs of curd until they are sufficiently dry. During this process, a continues to develop and the proteins probably undergo some change chemical constitution since their physical characteristics change. Sal now added which hastens the removal of the last of the whey, depresses action of the acid-forming organisms as well as the putrefactive ones. d is gently pressed, wrapped in cheese cloth and pressed again. It must

During ripening the enzymes present in the milk as well as those formed the microorganisms catalyze the transformation of many of the commods in the curd into other products. Sometimes an enzyme preparation deed at the time of salting. Ripening may be carried out at a temperature of 40° to 65° F, usually about 55° F; the lower the temperature the wer the ripening. A short time after ripening begins, the green cheeses dipped in paraffin so that the loss of moisture can for the most part be vented.

heddar-type cheese and all hard cheeses have relatively long keeping es. The cheese ripens uniformly through the mass and is not subject to growth of organisms on the surface until after it is cut. Often these eses are made in large 70 lb molds.

oft cheeses are made in small sizes, so that the microorganisms which elop on the surface can penetrate and the flavors developed will pass ough the cheese.

ening

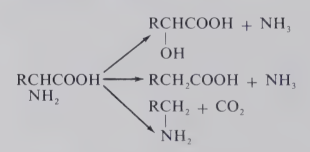
the changes which occur on ripening are numerous. Many investigations e attempted to trace the changes in chemical composition and the microanisms and enzymes responsible. Numerous reports continue to appear the literature as the attempt to understand and control the process, so ortant to the economic success of cheese making, persists. Only in very eral terms can we describe the changes that occur on ripening. Since the anisms in some green cheese are quite different from those in others, particular reactions will vary. In general, there are three types of the across in all cheese plus some others: (1) hydrolysis of protein under the across in a mine anide and forthy acids under the influence of specific and forthy acids under the influence of specific and forthy acids under the influence of specific and specific and forthy acids under the influence of specific and specific and forthy acids under the influence of specific and specific and forthy acids under the influence of specific and spe

(3) changes in amino acids and fatty acids under the influence of speenzymes to form flavorful compounds.

The hydrolysis of proteins results not only in change in texture but change in flavor. The longer the ripening, the more extensive the nges which occur. Large protein molecules are fragmented into smaller luble proteoses, into soluble peptones, into low molecular weight peps and amino acids. The enzymes which catalyze these changes include n and pepsin introduced in the rennet extract, as well as those synated by the microorganisms in the starter by which the milk was soured on the air of the cheese factory. Sometimes enzymes are introduced at time of salting. As the molecular weights of the proteins decrease, the

physical properties of the cheese changes. If most of the proteins are drolyzed to relatively low molecular weight soluble compounds, the chewill get very soft and creamy. This occurs as Limburger ripens. Even the fragmentation of the protein is not so extensive, the physical perties of the cheese change. The ability of ripened Cheddar cheese to readily and blend with liquids reflects a change in the protein. The molecular weight products of protein hydrolysis are flavorful. Some tones have a bitter taste but others are meaty. The amino acids for the part have relatively mild flavors. Some are sweet, some brothlike, others slightly bitter. All contribute to the blended flavor of the cheese.

- (2) The action of lipases on lipids, particularly fats, has a marked e on flavor. Butterfat contains a much higher percentage of low molec weight fatty acids which are soluble in water and volatile and conseque flavorful, than any other food fat. Hydrolysis of the fat liberates t acids and adds sharpness and their particularly distinctive flavor to cheese. Butyric acid has a very strong, rancid odor which is objection but small amounts blended with other substances account for the odo some cheeses.
- (3) Amino acids and fatty acids also undergo change which results the formation of some of the compounds most important for flavor. Nerous enzymes catalyze the deamination of amino acids with the format of ammonia and either hydroxy acids or simple acids. Both the ammonia and the acids are flavorful. Decarboxylation of amino acids results in formation of amines, some of which have very penetrating odors.



Changes in fatty acids can result in the formation of shorter acids eithrough beta oxidation of the fatty acid or through oxidation of unstrated acids at the double bond. Oxidations along the chain can result the formation of many molecules of low molecular weight acids or of tones and aldehydes. The high molecular weight acids, aldehydes, and tones are not water soluble nor volatile and consequently do not postlavor, but the low molecular weight ones are capable of stimulating eithe taste buds or the olfactory endings or both. For example, lauric a

indergo many changes depending on the enzymes present.

$$C_{11}H_{23}COOH$$
 COOHC₈H₁₆COOH + CH₃COOH

e oleic can be oxidized at the double bond as well:

$$CH_3(CH_2)_7CHO + CHO(CH_2)_7COOH$$
 $CH_3(CH_2)_7CHO + CHO(CH_2)_7COOH$
 $CH_3(CH_2)_7CHO + CHO(CH_2)_7COOH$

oquefort-type cheese the flavor depends to a great extent, although not

elv, on the presence of methyl ketones and butyric acid.

umerous other changes are possible and occur in some cheese. It has reported, for example, that the lactic acid formed during curd precipion disappears rapidly on ripening of some cheese while other acids ap-. Any lactose which may still be present in the green cheese also

opears.

ne organisms introduced from the air of the cheese factory have been tioned several times and since the character of the final ripened cheese ands so much on their activity, a little more should be said about them. en a large number of a certain strain of microorganisms develops in one n or building, the air and dust of that place contain these living nisms sometimes for years. If they are provided with a nutrient um in which to grow, the descendants flourish and in turn populate ir and dust with living members. Thus a wine factory where yeast entations are carried out year after year contains many strains of yeast. particularly those established in that place. A kitchen in which milk never soured will be practically free of acid-forming bacteria and if is left open to the air, it will putrify before it sours. But in a kitchen re milk has frequently soured, the air is so laden with these bacteria on exposure to the air milk sours in a few hours.

s the art of cheese making developed through the centuries, cheese I one area tended to be different from that of another even when inrents and methods were identical because the microorganisms estabd n the different factories were not identical. Roquefort cheese was prepared at Roquefort, France by ageing the cheese in caves. Until biology became a science, this cheese was only prepared in this town. at is possible to culture the mold which traditionally has been used

he ripening and add it to the curd.

the United States many of the cheeses developed in the Old World,

particularly in Europe, are successfully copied. By duplicating the ingre ents and the methods, and by adding the correct microorganism chee with very similar, if not identical, characteristics are produced.

Process Cheese

Processed cheese has become very popular in the United States because of its mild, standardized flavor and soft texture. It is produced by blenting aged cheese with green, young cheese in the presence of an emulsificand some water. The food sold as "processed" or "process" cheese has moisture content not more than 1 per cent greater than the average of the cheese used or in no case more than 43 per cent. That sold as "cheese food can have a moisture content up to 44 per cent while that sold as "cheese spread" can have a moisture content up to 60 per cent.

The expectation of the American consumer for uniformity in flavor doubtless one of the factors which has led to the popularity of these chee mixtures. Aged cheese from a single factory is usually *similar* in flavor are texture but not completely uniform, while that from two factories which use approximately the same techniques may differ considerably. A chee company catering to a nationwide clientele, is able to sell a product with the same flavor and texture in all parts of the country and at all times the year only by blending. Mild flavors in cheese products are likewise domanded by a large section of the American public. These mild flavors can be produced by blending a sharp aged cheese with green, young cheese with the curd.

The emulsifying agents used in the blending are salts: frequently sodiu or potassium phosphates such as NaH_2PO_4 , Na_2HPO_4 , Na_3PO_4 , $NaPO_4$, $Na_4P_2O_7$, $Na_2H_2P_2O_7$; potassium, calcium, or sodium citrate, $Na_3C_6H_5O_8$ sodium tartrate, $Na_2C_4H_4O_4$; or sodium potassium tartrate.

Processed cheese is produced by grinding aged cheese, warming, armixing thoroughly with young cheese, water, and emulsifier. Other foosuch as pimientos, olives, etc. may be added. Sometimes blends are man factured which contain two or more aged cheeses. An example is a chee blend which contains, Cheddar, Swiss, and Limburger.

Cheese *foods* are prepared by blending aged cheese, young cheese or cur with milk or milk products to form a smooth soft product. Cheese *spread* often have gums or gelatin added to promote smoothness and bind the spread together. Some cheese spreads contain no aged cheese at all, be only the curd. They have sugar, vinegar, and other flavoring substance added, but the "cheese" flavor is very mild. Cheese spreads are so soft the they are packaged in jars.

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CHAPTER NINE

Cereals and Their Use

Long before the beginning of the historic period, man learned cereals. Undoubtedly when primitive man was a food gatherer, he covered that the seeds of many grasses are valuable because the under proper conditions of dryness, be stored for long periods. La learned to grow grains in a little patch of cleared ground and assure self of food through the winter months. In the Western hemisphere or corn to Americans, was the cereal domesticated by the early In When Columbus discovered America, the inhabitants were growin most regions of the continent. Soon afterward maize was introduce Europe, and the grains of the Old World were brought to America early colonists. In Asia Minor and Asia wheat and rice were the moportant early cereals developed. Since the first wheat was grown in historic times and since before the time of recorded history many coccurred in the grain either through purposeful or chance breeding, cists as yet do not know what wild seeds were the parents of today's s

THE CEREALS

The cereals comprise a group of plants from the grass family, Gran whose seeds are valuable for food either for man or domestic are Wheat, barley, corn or maize, oats, rice, rye, and sorghum belong group; and although buckwheat is not a member of the family Gran it is usually classed with the true cereals. In some parts of the wor cereal is more important in man's diet than another. Usually the difference on the requirements of the particular cereal for soil, moisture length of time to maturity. Diet patterns are the result of centucustom where the availability of a particular food has established it. Thus rice, which requires warm, humid conditions and a long group.

cobetween 120 to 185 days from seeding to maturity), is eaten widely emhabitants of almost all the tropical regions of the world. Rve. if fourishes in poor soil and in cool climates, can be grown in northern ries and is relished by the inhabitants of northern Europe.

cereals give relatively high yields with a small amount of labor. Untedly this also was a factor in their early establishment as a cultivated

cture of the Grain

though the seed (grain) of each of the cereals differs from the others nere are likewise differences between subspecies like popcorn, sweet flint, and dent corn, all of these seeds are closely related botanically, detailed descriptions of each grain can be found in many textbooks of y, a generalized description will be given here.

e Primary Unit: The Cell. Like all living things, the primary unit in is the cell. Some descriptions of grains and other plant products by 1sts, home economists, or food technologists seemingly ignore the fact tells are present and that any cell is a complex mixture which always number of proteins, salts, many organic compounds in small amounts as the vitamins, molecules formed during synthesis or degradation, sually at least one form of carbohydrate, as well as lipids. Descripwhich ignore this fact are so oversimplified that they are highly misgrated the cells in cereals are living cells until milling occurs. Respiration wes during storage, although if the moisture content is low or if the erature is very low the rate may be very slow.

to of the Seed. The grain or seed of a cereal is in general composed of main parts: (1) the embryo or germ from which the root and leaf of w plant are formed when it sprouts; (2) the endosperm, the storage on of the seed which supplies the sprouting embryo with food in the before the root and leaf begin to function; and (3) the bran, which the covering or protecting layers. The three main portions of the seed adily be seen with the naked eye. The embryo is small and is attached base of the seed, while the endosperm makes up the major portion of d. The branny coats are composed of a variable number of layers of Si ce the functions of each of these three parts are quite different in

structures are readily differentiated in a wheat kernel. The bran is to possess six layers although botanically the aleurone layer, the lost one, is classed as the outer layer of the endosperm. However, in the aleurone layer separates with the bran. The peculiar property of e s of the branny layers, particularly after the wheat is tempered by

the controlled addition of water several hours or more before mill lows the ready separation of bran from other parts of the wheat se bran layers also have a very low density which assists in their sep from pieces of endosperm during milling.

Composition of Seed Parts. Chemically the bran is very different for rest of the seed. The bran has an unusually high per cent of crude file ash and a fair amount of crude fat. The crude fiber, it will be remer is composed of those organic molecules which are not soluble in dilute or dilute alkali (see p. 107) and includes for the most part the ce hemicelluloses, and some lignins. These substances make up the cel encrust the cell walls, or exist in the middle lamella between the cells.

The embryo (germ), like the bran, has distinct properties which is possible to remove most of this part during milling and which depend chemical composition. The germ is high in lipids and rather high is nitrogen and ash. During milling, the crushing and shearing action rollers squeezes out some of the lipids and causes a flat adherent fit the germ cells to form. The high lipid content causes the density of the to be low, and it is therefore readily separated from the endosperm parand the fine flour by shaking and bolting.

The endosperm is made up of cells containing large quantities of granules and cytoplasm, whose protein content varies with the loca the cell in the endosperm. Those cells close to the bran possess mortein and less starch than the cells in the center of the endosperm. The teins are most important in determining baking quality of the flour.

Storage of Grain

Easy storage of grains for relatively long periods makes them partivaluable to man during winter and has no doubt been an important in the prominent role that cereal grains have played in his history. proper conditions of dryness and temperature grains can be stored for periods with little or no change in either their fertility or milling and qualities. However, there are some problems. High humidity and higher peratures lead to mold growth, and infestation with insects is difficulties avoid so that long periods of storage are particularly hazardous.

Seeds are composed of living cells and during storage respiration tinues with the utilization of oxygen and the formation of carbon of and heat. Oxygen is readily available in the air held between the Wheat in bulk is one-third air. At low moisture levels, the rate of ration is slow; but as the amount of moisture in grains increases, so do rate of respiration. All grains have a critical level of moisture above respiration increases rapidly, causing heating of the grain and constitutions.

e. It has been demonstrated that the critical moisture level, about 14 per cent in wheat, is the level at which molds present on the grain the bran begin to grow. Their respiration is added to that of the ind gives an accelerated production of carbon dioxide and heat. Alnuthe amount of heat produced is sufficient to raise the temperature of ain, it is small and difficult to measure. However, carbon dioxide is ely easy to measure, and since it parallels heat production, it serves onvenient laboratory method for estimating heating.

ularly high. This can occur in grain where the average moisture conquite low when the grain is stored in large bulk and when irregular of it occurs. Moisture diffuses from warm areas to cool and the ure content in a cool area may rise to a point where mold growth grain is a poor conductor of heat, but as the mold flourishes, the recomes warmer and more favorable for this fungus. However, the eratures may become so high that the enzymes of the mold will be inted and the mold killed. This biological heating may be followed econd period of nonbiological heating in which the temperature may the ignition point. The whole process is termed "bin burning" and not or not fire results, there is marked damage to the grain in that on.

ects also contribute heat to grain since they have relatively high atory rates. The presence of insects in stored grain is always underbecause of the damage which occurs to the grain. Insect infestation y difficult to prevent and is one of the great problems of grain

e molds grow on grains, they invade the seed and by their enzymes ce compounds useful to their own development and life. These enaffect lipids, proteins, and carbohydrates, and serious changes in mposition of the seed occur. The germ may die and the milling and gualities are markedly affected.

the moisture content of the grain is of such critical importance for ping quality, grain is often dried before storage at the elevator. Drysto be carefully controlled so that sufficient time is allowed for diffusive to the surface of the seed and so that damage to the grain of occur.

AT AND WHEAT FLOUR

or pare a wheat flour, the miller faces the problem of separating the of the wheat so that they can be blended into a flour with a chemical issuon offering desirable baking characteristics.

Milling

Milling is a complicated process in which grain is ground in suc steps that gradually separate portions of it. Thus the bran is broken of flattened, the germ is pressed into a flake, and the endosperm is pow Clean wheat is "tempered" before grinding by treating it with water s the bran will be tough and readily separated from the endosperm. Th pered wheat is crushed between corrogated rollers called break roll first break rolls are set relatively far apart and grind the wheat I while successive breaks yield finer and finer products. The first break arated by sieving or bolting into very fine particles (flour), interm particles (middlings), and coarse particles (chop, or stock). The ch stock is then sent to the second break rolls. This process may co through 5 or 6 breaks. The chop contains pieces of endosperm and and the chop from the last break is principally bran. The middlings co endosperm, bran, and the germ. The middlings are classified and so the bran removed and sent to the reduction rollers. These are smooth ers, but like the break rolls they are graduated so that successive redu becomes finer and finer. After each reduction, sifters separate the middlings, and chop. This process is continued until most of the endo has been removed as flour and most of the bran has been separated sifters. The remnant consists of fine middlings, bran, and a little ger is used for animal feed.

Types of Flours Produced. The flour streams from the various rolled named for the roller—"first break stream," etc. They vary in che composition because they vary in the amount of bran and germ which contain as well as in the portion of the endosperm which has been released. Many different flour streams are produced, especially in large mills, streams are combined and blended to form flours with particular becharacteristics. Straight flour is a combination of all the flour stream is seldom produced. Patent flours are flours from the more refined stand vary considerably in the percentage of the total flour representations.

Family patent: 70–75 per cent total flour Short patent: 75–80 per cent total flour Long or standard patent: 90–95 per cent total flour

The flour remaining after the patent is removed is called *clear flour* too may be separated into different grades. Clear flours are used in bination with rye for dog biscuits and in other products where high q is not essential. The flour from the last reduction is called *red dog* be

sually sold as animal feed.

lation to Dough. Milling does not involve any chemical procedures. It is a physical separation of parts of the wheat seed, but it does affect reformance of the flour in the dough stage. Starch is held in granules cells of the wheat seed, and on milling the cells are ruptured and the granules escape. The grinding also damages some of the granules so starch flows out of the granule case when the flour is made into dough, the available for hydrolysis through the action of α -amylase. In flours the amount of damaged granules is around 4 per cent. These ged granules are called "ghosts" and the level at which they occur is retain in determining the amount of α -amylase activity at room temperature in the dough.

analyses of the various mill products must be considered as only repative since wheat, like all plant products, shows variation in comon with variety, climatic conditions, and soil fertility. Milling does impletely separate bran, germ, and endosperm. Thus, red dog conconsiderable amounts of bran and germ particles, and the analyses rehis in the relatively high crude fat and crude fiber concentrations, able 9.1.

Proteins

· flour proteins are of most importance as far as baking quality of is concerned. The large molecules of proteins are readily modified id treatment, both physical and chemical, and consequently their

TABLE 9.1. CHEMICAL COMPOSITION OF CERTAIN MILL STREAMS AND BY-PRODUCTS OBTAINED IN WHEAT MILLING.

roduct	Moisture (%)	Total Nitrogen (%)	Fat (%)	Fiber (%)	Pento- sans (%)	Ash (%)	Total Sugars	P ₂ O ₅
	10.3	2.05	2.1		5.1	1.73	2.6	() ×5
'et i Flour	11.5	1.82	1.0	0.2	2.9	0.40	1.3	0.22
ar Hour	11.0	2.13	1.7	().2	3.6	0.81	18	() 44
'Tear Flour	10,4	2.33	2.0	0.3	3.4	1.34	21	() ~()
	9.2	2.87	5.4	2.4	8.4	3 15	64	163
	8.8	2.33	4.1	10.8	251	6.38	5.4	3 13
	8.9	2.47	5.2	8.4	16.3	4.10	6.0	2.23
	8.5	4.84	11.9	1.8	6.2	4.80	15.1	2.73

me toon of data of C. H. Bailey by Geddes, W. F., "XXIV, Cereal Grains" in Jacobs, M. B., ed., "mistraind Technology of Lood and Lood Products," Interscience Publisher Visit N. Y.

physical properties are changed by these treatments. Some of these fications occur in doughs and batters.

At the beginning of the century Osborne studied the proteins of ber of cereals and separated them on the basis of solubility (i.e. ability to form colloidal dispersions in certain water solutions) in categories: (1) the proteins soluble in 70 per cent alcohol, calle lamines, (2) the heat-coagulable proteins soluble in water, called alb (3) the proteins soluble in neutral salt solutions, called globulins. proteins soluble in dilute acid and alkali, called glutelins, and (5) defined fraction called proteose. In the many years since Osborne's p work many proteins have been fractionated on the basis of their solu However, now that the ultracentrifuge, electrophoresis, and di make it possible to demonstrate whether molecules in a given fraction the same size or range of sizes, it has been found that fractions bas solubilities do not give single proteins but a mixture of them. Whea globulin contains at least three proteins while the albumin fraction h Keeping in mind the limitations of the methods of separation bas solubility, we can nevertheless learn a little about the proteins of by considering the results of this method. In all cereals the chief pr are prolamines and glutelins. In corn or maize the prolamine (zein) fr is the most abundant protein, while the amounts of glutelins are lo oats and rice it is the glutelins (called avenin in oats and oryzer rice) which are most abundant. In wheat, rye, and barley the propo of prolamines and glutelins are intermediate. In the wheat endospe prolamine (called gliadin) and a glutelin (called glutenin) are presi approximately the same concentrations; in the bran a prolamin is abundant with fair amounts of an albumin and globulin. Althou cereals are more or less similar in their protein content, the unique ence of glutenin and gliadin in the wheat endosperm is important baking operation. In the presence of water and with mechanical agi these protein fractions form a tough, elastic complex termed gluten is capable of retaining gases and by so doing makes a leavened pr possible. Cereals other than wheat cannot form a large light loaf gluten is not developed.

Gluten. It must be remembered that the terms gliadin and glutenin dindicate homogeneous proteins but rather protein fractions. Likewisterm gluten does not apply to a group of identical molecules. Glute be readily prepared by adding 60 to 65 per cent water to a hard flour, allowing the dough to stand approximately 30 minutes, and washing out the starch granules and soluble compounds under a strewater. A tough, elastic, gummy product is obtained which consists of

mately two-thirds water and one-third protein. There are also small ants of lipids, starch, and ash. Sullivan²¹ gives a typical analysis of gluten: protein, 85 per cent; lipid, 8.3 per cent; starch, 6.0 per cent; starch, 6.0 per cent; our and the method of handling. The lipid is held strongly to the mand cannot be extracted with ethyl or petroleum ether.

Ivan also reviews briefly the evidence which Hess has presented to that gluten does not occur as such in the endosperm but is formed gh mechanical treatment of the flour in the presence of water. Protein ited from flour and freed of the starch granules by methods which it use water shows a swell in contact with water of 25 per cent. Gluten in contact with water 200 per cent. The swelled particles show diffiraction patterns with X rays. These observations indicate they be different proteins.

anature of the chemical reaction by which gluten is formed has cons cereal chemists for many years. Although the product is always gummy, elastic, and tough, the degree to which these properties are sped varies from flour to flour. Both the "machining qualities," the which a dough can be handled by a mixing machine, and the baking ities depend on a nice balance between stickiness, extensibility, ness, and tenderness. As vet there is no chemical explanation for the nces in gluten produced from different flours and the only way in the quality of the flour for machining and baking can be estimated arrying through the operation. It was once believed that differences · relative proportions of glutenin and gliadin might explain the nces in the gluten. But flours that differ markedly in their baking rties show little or no difference in the classical fractions of glutenin radin. Other investigators once hoped that when methods for the deration of the amino acid content of proteins had been perfected, this account for the difference in quality. Today those methods are well ped: and although the proteins of flour have an unusual distribution ino acids (high in the amides of aspartic and glutamic acid, parly glutamic, and relatively high in leucine and proline), there are no and variations in the amino acid content of glutens of widely differperties.

as been traditional to consider the development of gluten in a flour er mixture as the reaction of two protein components, gliadin and

Many types of studies have shown these to be mixtures rather mule proteins. Attempts to disperse gluten, gliadin, and glutenin ous solutions have resulted in fractions which are heterogeneous, phoresis, diffusion, and sedimentation by the ultracentrifuge indi-

cate numerous components present in each of these proteins. The p of the structural composition of gluten is far from solved; and utime when it is solved, it is convenient to use the old names—gliadin, and glutenin.

Strength of Gluten. The properties of gluten formed from differen or with variations in procedure, differ. The stickiness, toughness, elabrittleness, and coherence are all part of the "quality" of the gluten f A gluten which is elastic but still fairly tough is called "strong" wh which is quite sticky and not very elastic, which spreads when a placed on a plate, is called "weak." Many factors influence the strea gluten formed from a single sample of flour, and account for the ence in glutens formed from different samples. The effect of all factors to completely understood.

In general, hard wheat gives gluten of good strength while soft forms gluten of low strength. In the past it has been suspected the total protein content of a flour is a measure of gluten strength. This true since flours with the same percentage of protein may have strengths which are widely different. It is true, however, that hard flours which form the strongest gluten have the highest percentage tein. Protein content has not explained the difference in gluten stand cannot be used as a means of evaluating the gluten strength of Unfortunately, the knowledge of the chemistry of gluten has not ad sufficiently to allow measurement of differences in gluten strength by ical analysis.

The problem of gluten development is further complicated by that hydration of the protein occurs as the gluten develops. Many p become hydrated and hold water tenaciously. The protein swells a water is not readily removed by ordinary drying methods. Gluten this same phenomenon. Hydration capacity is influenced by many such as pH, total ion concentration, and presence of soluble mosuch as sucrose. The properties of a dough or batter also vary with p concentration, and sugar concentration; and doubtless the effect of conditions on gluten hydration is one of the factors which produce in properties. The role of gluten in baked products will be discussed doughs and batters.

Lipids and Carbohydrates

The *lipids* of wheat are concentrated in the germ, so that althous wheat seed contains only about 2 per cent lipid, the germ contains of 12 per cent. This does not mean, however, that all the lipids are the germ. Flour always contains 1 or 2 per cent lipid. The amount

mate and soil under which it is grown. The lipids present include alglycerides as well as phospholipids and sterols. Wheat germ oil is a nts of fat soluble vitamins and pigments separate with the ether extended fat."

e principal carbohydrate of all cereal seeds is starch but there are is small quantities of others present. Dextrins are present in small nts, particularly in flours since they are formed to a limited extent the starch on milling. The water soluble carbohydrate is sucrose and fore abundant in the germ than in the endosperm or bran. Some cellupresent in all parts of the seeds since it is the chief component of il walls. There are also small amounts of lignins and pentosans in the hemicelluloses and pectic substances mainly in the bran. Water exof either the whole grain or a flour yield gums. In wheat gum there oth pentosans and hexosans. A pentosan composed entirely of xylose cen isolated from durum wheat flour, and one composed of arabinose ylose has been isolated from spring, winter, and durum wheats. 84.)

nins

vitamin content of cereal grains has been studied extensively. All the grains contain important quantities of the vitamins of the B complex; here the use of cereals is sizeable, they contribute significant quantithe adequacy of the diet. Ascorbic acid is completely lacking in the riour. Vitamins A and D are absent, but yellow corn derives its yellow corn carotenoids. The pigments in yellow corn are principally xanthin with small amounts of α - and β -carotenes, which are presof vitamin A. Wheat also contains carotenoids, principally xanthowhich has no vitamin A activity in man or other animals. (See Caros, p. 228). The germs of various cereal grains contain vitamins E, the serols, which are pressed out or extracted with the lipid fraction. germ is particularly rich in the tocopherols.

min, riboflavin, niacin, and vitamin B_a are present in fair amounts in the cereals while pantothenic acid and folic acid are in smaller. The quantity of these vitamins present in any specific sample of four shows the same variation of values which is found in any plant differing with the conditions of climate and soil under which it man and with the variety and species. Thiamin varies from $3 \mu g$ per g herice to 90 μg per g of oats. In 99 samples of wheats the thiamin

ried from 3.2 µg per g to 7.7 µg per g.

OF WILLIAM							
Product	Mill Yield (per cent)	Thiamin µg per g	Riboflavin µg per g	Niacin µg per g	Panto- thenic µg per g		
Patent Flour	63.0	0.68	0.34	12	5.7		
First Clear Flour	7.0	3.0	0.62	26	9.6		
Second Clear Flour	4.5	12.37	1.85	83	12.8		
Red Dog Flour	4.0	29.66	3.80	120			
Germ	0.2	22.93		68	15.3		
Shorts	12.3	17.40	2.80	159			
Bran	9.0	9.37	2.80	330	Miller		
Wheat	100.0	5.02	1.00	70	12.2		

TABLE 9.2. DISTRIBUTION OF VITAMINS IN PRODUCTS
OF WHEAT MILLING*

The riboflavin content of whole grain is lower than that of thial does not show quite as wide variation for the different cereals. I tween 1.1 and 1.6 μ g per g.

The niacin values for grains show some disagreement becamethod influences the data obtained. With alkaline extraction, the are higher. In wheats the quantity ranges from 47 to 106 μ g per grains are in the same range, although rye and oats are lower.

The distribution of some of the B complex vitamins in different of the seed or in different flour streams has received considerable as in recent years with the growth of interest in food fortification. In the branny layers are richer in the B complex vitamins. Thus the distributed in approximately these amounts: bran, 61 per cent; end 24 per cent; germ, 15 per cent. While all flour contains small per of germ and bran, highly milled flours with good baking quality are most part from the endosperm. Table 9.2 gives the amounts of B ovitamins in various mill products.

Enrichment of Flour

It will be noticed from Table 9.2 that the amount of the B comp mins in flour is considerably lower than that in the whole wheat g the diet of many individuals the consumption of white flour product sents about one-fourth of the total calories. If the rest of the die in B complex vitamins, the small amount in the white flour does not But for many Americans this is not true. The diet is low in the vitamins as well as in iron and calcium. Iron is likewise rather a in the branny layers of the wheat kernel but largely removed on For many years nutritionists and doctors have recommended the

^{*}Part of a table in Geddes, W. F., "Cereal Grains," loc. cit., p. 1100.

TABLE 9.3. STANDARDS FOR ENRICHMENT OF FLOUR EFFECTIVE OCTOBER 1, 1943

	Conten Min.	t per lb
Fequired Ingredients		
Thiamin, mg Riboflavin, mg Niacin or Niacin Amide, mg Iron, mg Optional Ingredients:	2.0 1.2 16.0 13.0	2.5 1.5 20.0 16.5
Calcium, mg Vitamin D, U.S.P. Units Wheat Germ	500 250 Not more	625 1000 e than 5 per cent

wheat. But the recommendation has been largely unheeded. Whole coes not produce as large or light a loaf of bread or as light a cake white flour. In recent years many have advocated the enrichment of flour to its original level with thiamin, niacin, riboflavin, and iron. It a new Food, Drug, and Cosmetic Act, which became effective on per 1, 1943, prescribed the amounts of these compounds which must orporated in enriched flour and the amounts of calcium, vitamin D, heat germ which are allowed. See Table 9.3. Some nutritionists have add the addition of calcium and vitamin D, since they are not nat-present at those levels.

re has also been interest in the fortification of bread with lysine. The ceins of wheat flour contain little lysine; and when they are the sole of amino acids in the diet, an animal soon develops deficiency onto the low priced diets for man often contain bread or other wheat it is in abundance since they supply sufficient calories cheaply. In these here are not large amounts of other proteins to make up for the lack he in wheat and mild deficiencies can occur. Two methods are now or this extra fortification: (1) addition of free lysine to the dough tuse of skimmed milk powder. Addition of lysine does not alter the structure, but incorporation of much skimmed milk powder pro-

In parts of the world the staple food for man is either rice or either of these cereals have complete proteins. Rice is low in both and threonine, while corn is low in lysine and tryptophan. Rosenta shown that growth of white rats is improved by supplementated the of cooked rice (90 per cent) with lysine up to 0.1 per cent

but that improvement falls off at higher levels of lysine. On 90 per corn meal diet rats grow slowly, but this growth can be improved by addition of up to 0.05 per cent lysine hydrochloride.

The *mineral* elements are present in the plant cells sometimes as and sometimes as part of organic molecules. We know, for example, phosphates occur as hydrogen phosphate, HPO₄=, and dihydrogen p phate ions, H₂PO₄-, but they also occur as the acid portion of many ganic esters such as glucose-1-phosphate, lecithins, phosphopyruvic a and many others. The mineral elements are determined from the ash of ural products where the organic molecules have been destroyed and the noncarbonaceous elements remain. Cereals contain a large numbe elements, although many of them are present in trace amounts. Calcimagnesium, potassium, sodium, iron, phosphorus, chlorine, sulfur are present, and in the cereal seeds that contain hulls (covered grain) that are sizeable amounts of silicon. Traces of aluminum, arsenic, boron,



FIGURE 9.1. MRS. CATHERINE CLARK OF BROWNBERRY OVENS IN HER EXPERIME KITCHEN. A high protein bread supplemented with lysine is one of her specialties.

Courtesy of Brownberry 0

e. copper, cobalt, iodine, lithium, tin, selenium, titanium, and zine are ent. As a group the cereals are not important sources of the mineral ents; and since the bran is richer in most of these elements than the sperm, highly milled white flour and products baked from it are poor ses of minerals. Calcium, the mineral most often poorly supplied in is diet, occurs to the extent of only 0.15 mg per g in corn, 0.5 mg per g heat, and 1 mg per g in oats.

EATS AND THEIR BAKING QUALITIES

heats grown under different conditions of soil and climate and wheats flerent varieties have different baking qualities. Some wheats yield a which will produce a loaf of bread of great height and good texture, rs will form a small loaf of poor texture. Likewise in batter products ightness and grain depend on the flour used.

bday in the United States technology of foods has progressed to the twhere the consumer expects a uniform product and will not be satisatith less. In order for the baker to turn out a loaf or cake exactly like one produced last week, it is necessary to understand the factors which unt for a good loaf or a poor, a good cake or a poor one. During the fitty years much research has been done on flour products. But so comis a loaf of bread or a cake in its chemical and physical-chemical reaships that all of the questions have not been answered. The problems p around three main areas: gas production, gas retention, and tender. In the yeast-leavened doughs of bread, rolls, coffee cake, and crackers roblem is a little different from that of the batters of cakes, cookies, ns, and biscuits. Each will be briefly discussed under Doughs and then r Batters.

ghs

s Production in Yeast-Leavened Doughs. The common agent for the action of gas in doughs is a selected strain of the yeast. Saccharomyces side. The yeast through its life process forms carbon dioxide as an tory product and in so doing leavens the dough. The yeast also grows 2 the fermentation period and produces numerous offspring which is excrete carbon dioxide. The amount of gas formed depends on the reast and on the number of organisms present during fermentations strains of yeast produce more gas in a given time than others. The product is also important because carbon dioxide is only one 11 compounds excreted, some of which have flavors.

technology is not as advanced as in the United States, the yeast used

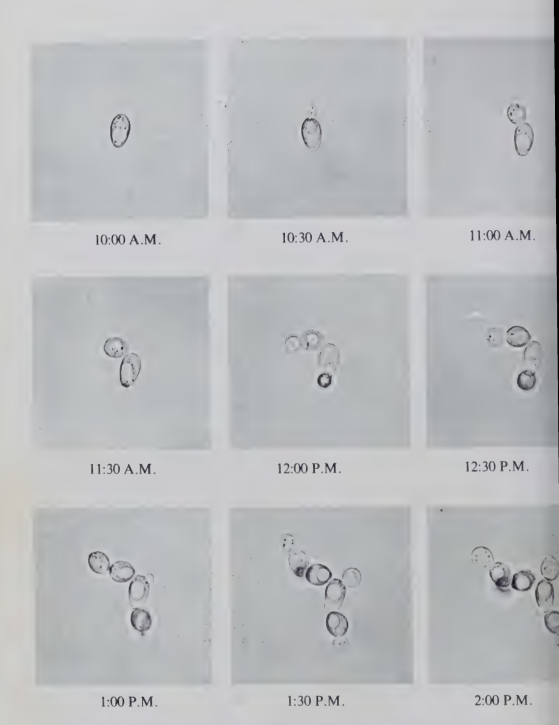


FIGURE 9.2. GROWTH OF YEAST CELLS. Photomicrographs taken at half-hour int Magnification × 1000. Courtesy of Fleischmann Laboratories. Standard Brane

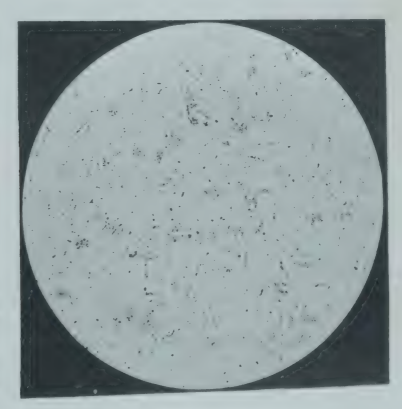
by bakers is a "starter," saved from the last baking. The "starter" is a mixture not only of yeasts but also of bacteria and the rate of grow the various organisms depends on the nutrients present as well as the perature at which they are grown. Contamination with "wild" yeasts bacteria readily occurs in kitchen or bakery. A baker might have

es with the organisms.

took many years for the companies which produce yeast to learn how dequately protect their cultures and encourage the growth of selected its of organisms as well as how to package them so that they reach the immediate and highly viable. Much effort is devoted to maining a uniform strain. Today uniform yeast cultures are available either impressed, dried, or powdered yeast.

"sour dough" products bacteria which produce carbon dioxide and mic acids are used for either part or all of the leavening action. Alagh these organisms were formerly preserved as starters, most bakers as use cultures or some source of a uniform culture like cultured butterinstead of depending on an unreliable and variable "starter." Some of organisms used are the acetobacters which form acetic acid. Strepto-us lactis, and the lactobaccilli, L. acidophilus, L. bulgaricus, and L. i, which form lactic acid.

he Nutritional Needs of Yeast. Since a yeast cell is a living organism, it numerous nutritional needs and it is only if these are met that it will a vigorously and produce a large quantity of carbon dioxide. Some a of easily available carbohydrate and a utilizable source of nitrogen, ium, and phosphate ions are important in the dough for rapid gas nation.



URE 9.3. COMMER-PE BREAD SOURING SM STREPTOCOCCUS 15. Magnification ×

nurtesy of Fleischmann aboratories. Standard Brands, Inc Yeast cells use a number of mono- and disaccharides. These sugars metabolized to ethyl alcohol and carbon dioxide. Glucose and fructose the monosaccharides likely to be in yeast-leavened doughs, while such and maltose are the disaccharides. Lactose may be present but is not by the yeast cells. The sugar in flour is mostly sucrose with only samounts of glucose and fructose. Most of the glucose and fructose is in wheat seed in the germ; and since this is chiefly removed in the milling, amount in the flour is low. Sucrose is present to the extent of 1 to 2 per in the flour, and a yeast, flour, and water mixture will ferment rapidly this is used up. Too much sucrose, however, will slow down the rate of mentation. If a very sweet dough is prepared by adding 10 per cent or not sucrose at once, the growth of the yeast and the formation of carbor oxide may be slow. Maltose is not present in flour to any extent; but we the dough is made, the amylases begin to hydrolyze starch and formaltose.

Amylase Content and Activity. The amylase (or diastase) content of dough has an effect on the rate of gas production by the yeast because the enzymes form sugars from starch. Two types of amylases occur in nat α -amylases which hydrolyze the internal 1-4 links in the amylose chain the amylopectin brush, and β -amylases which hydrolyze the 1-4 links ond from the end of the chain. Bernfeld² says that ungerminated cereals not possess α -amylases but that they are formed during germination malting. β -Amylases occur in both the ungerminated and the germinate cereals. α -Amylases through their hydrolytic action on internal 1-4 links applied produce dextrins of various sizes and then form smaller and small molecules. The final products are maltose and some glucose from amylobut there may be larger molecules from amylopectin since the 1-6 links not attacked. (See p. 76.)

 β -Amylases hydrolyze the 1–4 link second from the end of the chain a consequently rapidly form maltose. On hydrolysis the configuration of bon 1, the potential aldehyde, is changed from the alpha to beta configuration. Under optimum conditions β -amylase can catalyze the hydrolysis the entire amylose chain to β -maltose; but with amylopectin, the 1–6 liare not attacked and a terminal dextrin is formed.

 β -Amylase is found naturally in flour in small amounts and both α - β -amylases in greater amounts in malt extract, malted barley, or may wheat but not in yeast. Since the formation of maltose from starch is sential for the adequate growth of yeast in an unsweetened dough, the least of these enzymes in the dough is of paramount importance. They can added to the dough or flour either in the form of malt extract or as enzyme concentrate. Today most millers carefully regulate the amount

lases in the flour so that the flour will have proper gas production ities when it is inoculated with yeast. Some patent flours are low in lase activity, and this is rectified by the addition of malted wheat flour halted barley flour.

amylase activity is great enough, the production of maltose will keep out the demands of the speedily growing yeast and carbon dioxide torsion will be rapid. Where amylase activity is low, the addition of dexand soluble starch will increase gas formation. Preparations composed nese compounds do not improve gas formation except when amylase vity is low. Overgrinding has a similar effect in the presence of low lase activity by rupturing some of the starch granules and making them e readily hydrolyzed by that amylase which is present.

mylase activity has an optimum pH of 7 and, if the dough is at some or pH, then amylase activity is lower. The addition of milk to the gh raises the pH because of the presence in the milk of buffer salts. It consequently retards amylase activity. However, in the presence of salts such as calcium hydrogen phosphate or the acetic acid of vinethis retardation may be eliminated, and gas formation may even be inseed by the milk through the improved nutrition of the yeast.

turing baking the starch held in granules imbibes water, swells, and is tinized. The β -amylase from either malted barley or malted wheat is tivated at a higher temperature (approximately 175°F) than the temperature for the gelatinization of wheat starch (approx. 150°F). Consently, during the early stages of baking β -amylase affects the gelatinized α . The baked loaf achieves an internal temperature close to 212°F and his temperature enzymes are destroyed. See figure 9.4.

Talt, malted wheat, and malted barley are the sources of enzymes comily used. These products contain not only α -amylase but proteinases ell. Both types of enzymes exert a pronounced effect on the dough, and level at which they are added can be critical. In these products the lof enzyme activity is variable. Today purified enzyme extracts from a or bacteria are available and considerable investigation is being otted to their use in doughs. With more sources of enzymes in which the lase or proteinase levels are more accurately known, it will be possible introl dough performance more carefully.

Drugh conditioners" or "flour improvers" are on the market and conlinear and phosphate ions. Both of these ions are necessary for yeast th and gas formation but flour contains them in sufficient amounts mod leavening under normal conditions.

re ferments. These have been used for many years in bread making, but nt y their effectiveness in commercial baking has attracted attention. A

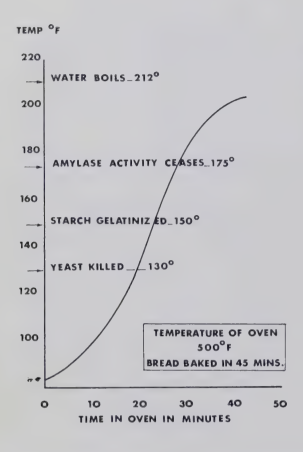


FIGURE 9.4. TEMPERATURE AT THE TER OF A 2 LB LOAF DURING BAKIN produced from Horder, T. J., Dodds, and Moran, T., "Bread," Constable don, 1954.

pre-ferment is a more or less fluid mixture to which yeast is added and which it grows rapidly. It differs from a sponge in the absence of or in a level of flour. Potato water and yeast or a liquid mixture of mashed be potato and flour are examples of pre-ferments which have been used in past. Johnson, Miller, Refai, and Miller¹⁰ point out that Parisian B was brought from Paris to Scotland in 1865. It was a pre-ferment mof malt, salt, and flour inoculated with yeast and was used not only source of yeast but also to improve the flavor of bread. While the yea developing in the pre-ferment, it forms not only alcohol and carbon dio but other compounds such as organic acids which affect flavor.

Johnson, Miller, Refai, and Miller have studied six pre-ferments w differ in carbohydrate used (either glucose or sucrose) and the presence absence of dry milk solids and flour. Their mixtures provide the nutri demanded by yeast for rapid growth. In all mixtures containing sucre the carbohydrate disappears immediately when it is mixed with yeast. Trapid hydrolysis of sucrose by yeast had been known before. There are marked differences in the six pre-ferments at the end of 5 or 6 hours. We milk solids are present, the rate of utilization of the carbohydrate slightly slower and because of the buffering action of the milk salts, pH does not fall as low. There is a slightly greater formation of carbohydrates.

de in the presence of milk solids. Production of organic acids is apamately the same in all pre-ferments, reaching a maximum in 3 hours then falling.

as Retention in Doughs. When flour is mixed with water, a change takes e in the physical-chemical properties of the gluten. It has been customto consider gluten the product of two protein fractions of wheat din and glutenin. (See p. 322.) It is in wheat flour and only in wheat r that this elastic, rubbery protein develops when it is stirred with er. The ability of a dough to retain gas depends directly on the unt and quality of the gluten developed. The volume or lightness of a at loaf is therefore directly dependent on the factors which contribute he development of gluten and which influence its elasticity. Doughs le of other cereals are not able to retain all of the gas generated and sequently are heavy and coarse textured. The factors of major importfor gluten development and strength in most doughs are: (1) menical action, (2) the amount of proteolytic enzymes, (3) the amount xidation of the flour either through aging or the addition of oxidizing its, (4) the addition of milk, inorganic ions, etc., and (5) the addition icrose and fat of importance in sweet rolls and coffee cake.

deither through mixing or knéading is important for the development te gluten. Elastic strands of rubbery gluten form as the flour is mixed water. Undermixing produces a dough in which there has not been cient gluten development to retain the gas well and the result is a loaf h has poor volume and is heavy. Overmixing or rough handling of the 2h before panning has been reported to decrease the volume of the loaf. In flour and water are mixed, an increase in plasticity occurs as the en is developed. But if mixing is continued, a decline in plasticity occurs.

and the dough finally becomes slack and sticky.

roteolytic Enzymes. These are a group of enzymes present in wheat and which catalyze the hydrolysis of proteins. They are also present in ed wheat flour, malted barley, malt extract, and yeast. Since the 19th of the gluten depends on the intact protein, any reaction which olyzes part of the protein reduces the amount of gluten. If too much of 19th proteolytic enzymes are present, too much hydrolysis occurs and the 19th becomes sticky, difficult to machine in the mixers, and yields bread of 19th volume. However, some protease activity is desirable since it improves 19th uten. Doughs of low proteolytic activity are called "bucky" because 19th uten is tough and inelastic. It does not machine well and it also protection is 19th uten in the 19th uten in the 29th uten. The complexity of the problem, "what is gluten." Is well demounted to 19th uten. The complexity of the problem, "what is gluten." Is well demounted to 19th uten.

onstrated here. The quantity of proteolytic enzymes must be balance that enough hydrolysis occurs to produce an elastic gluten but not so that the gluten is sticky. The addition of malt extract, malted wheat or malted barley must be limited to avoid an excess of proteolytic acti

Oxidation. Oxidation of flour likewise influences the buckiness or ness of the gluten developed and here, too, a nice balance must be ach It has been known for many years that storage of flour at moderate peratures for several months lightens the creamy color and improv baking quality. Eventually it was found that oxidizing agents can a this effect without the costly and hazardous period of storage. I United States the Pure Food, Drug, and Cosmetic Act of 1938 allow use of chlorine, chlorine dioxide, nitrogen oxides, nitrosyl chloride benzoyl peroxide. Nitrogen trichloride, "Agene," was used for many but on August 1, 1949, it was removed from the list of permissible a See p. 352. It was found that dogs, cats, and a number of other speci large amounts of flour bleached with Agene or bread made from that developed running fits and convulsions. Although no harmful effects found in man, the compound is no longer used. Chlorine, chlorine die and nitrosyl chloride bleach and mature the flour, improving the prop of the gluten formed in working the dough. Nitrogen dioxide and be peroxide do not affect the baking properties of the flour at the levels in milling but only exert a bleaching action.

Potassium bromate, KBrO₃, and potassium iodate, KIO₃, are added as dough conditioners. They are oxidizing agents and are believ have an effect similar to oxidizing agents added at the time of milling amount of potassium bromate which exerts this effect is surprisingly so the order of one or two thousandths of one per cent.

The effect of either oxidation or reduction on flour and dough has studied for many years, but the exact nature of the reaction is still in d If an oxidizing agent is added either to flour or to gluten, the streng the gluten is increased. If relatively large amounts are added, the glute comes tough with little elasticity. Reducing agents have the opposite of They cause a decrease in the strength of the gluten, making it more ex ble and sticky. When a reducing agent is added to gluten, a ball of it the ability to stand up and soon softens and flattens. Sullivan²¹ point that numerous factors influence the physical properties of gluten—fer tation, oxidation, the work of mixing, rounding, braking—and that these can be considered active in changing the bonding of the protein cule. She reviews some of the evidence which indicates that oxidation fects the sulfhydryl groups and increases the number of sulfur-sulfur li

$$2 - SH + (O) \rightarrow -S - S - + H,O$$

the molecule together more firmly and increase its strength. Overoxical flour can be used satisfactorily by increasing the working which it resome of the less stable bonds between polypeptide chains are damit and the molecule is rendered more extensible.

hether the effect of oxidation is on sulfhydryls or some other group. It important reaction in practical baking. The amount of oxidation is cal and although the amount of oxidizing agent needed is small, suffimust be present to yield gluten strong enough to retain the gas and but not so tough that it will be unable to stretch around the gas bles.

ther Factors. Several other ingredients may have an effect on the 19th of the gluten. Raw or pasteurized milk decreases the baking qualifor a flour unless the milk is first heated. It is believed that milk consome substances which increase the activity of the proteolytic enzymes consequently during the fermentation period foster the formation of a en which is too sticky. Heating the milk to 180°F for 30 minutes roys these unknown substances and the milk, then, has no detrimental it on gluten strength. In home baking it is always recommended that milk be scalded before use in a dough. Pasteurization also provides a od of heating but for a much shorter length of time or at a lower temture. The U. S. Public Health Service requires that pasteurized milk be at 143°F for 30 minutes or at 160°F for 15 seconds.

hen the fluid used in a dough is water, the softness or hardness of the er has an effect on the gluten. In general, calcium salts present in the water tend to increase the elasticity of the gluten. Sodium chloride vise affects the gluten. Acids also alter gluten strength. Vinegar is somes added to dough, but too much must be avoided so that the gas retenis not diminished. Fats in small amounts increase the ability of the zh to retain gas.

ne production of a uniform high quality loaf or roll is complicated by y factors. Much empirical work goes into the formulation of recipes the control of the quality of the ingredients. The complex problem of physical-chemical properties of a dough are still incompletely under-

ers

Production in Batters. The satisfactory production of baked goods butters involves the same type of problem as from doughs: adequate ormation and gas retention. In batters the leavening agent is either

air which is incorporated through beaten egg white or during cream it is a gas generated by chemical reaction in the batter from a powder or from ammonium bicarbonate. Often a combination of lea agents is used. Thus in the production of some butter cakes, creaming and fat introduces air and later in the mixing the addition of beate white introduces still more air. Baking powder is added which in the batter and at the oven temperature forms carbon dioxide. Steam als as an important leavener.

Baking powders are composed of sodium bicarbonate, NaHCO₃, a salt, and starch. Occasionally some other ingredient such as a small at of egg white is present as a drying agent. Starch helps keep the ingredry, prevents caking and standardizes the baking powder so that for a volume of dry baking powder the amount of gas formed is equal to to other baking powders. The chief difference is, therefore, in the compounds which produce hydrogen ions (hydronium ions) to with sodium bicarbonate in the wet batter and release carbon dioxide.

$$NaHCO_3 + H^+ \rightarrow Na^+ + H_2O + CO_2$$

Phosphate baking powders contain either primary calcium phosp Ca(H₂PO₄)₂ or disodium pyrophosphate, Na₂H₂P₂O₇. Those on the sumer market contain primary calcium phosphate, while some of the phate powders for bakers contain disodium pyrophosphate. Tartrate ing powders contain potassium acid tartrate (cream of tartar), KHC₄F as well as tartaric acid, H₂C₄H₄O₆, as a source of the hydrogen ion. S baking powders contain sodium aluminum sulfate (alum), NaAl(S The hydrated aluminum ion hydrolyzes on contact with water and for hydronium ion (hydrated hydrogen ion).

$$Al(H_2O)_6^{+++} + H_2O \Rightarrow H_3O^+ + Al(H_2O)_5(OH)^{++}$$

The phosphate and tartrate baking powders react readily at room term ture to form carbon dioxide. S.A.S. baking powder forms little carbon dioxide at room temperature, and it is only at oven temperature that leavening occurs. If much time elapses between mixing and bathen the slow acting S.A.S. baking powder is much better. Leavent room temperature is not effective unless baking follows immediately so the gas formed will not escape from the batter. *Double action* be powders that contain both sodium aluminum sulfate and primary caphosphate are available on the consumer market. An S.A.S. powder the disadvantage of forming sodium sulfate which has a distinct flavor and is therefore seldom used alone.

mmonium bicarbonate or ammonium carbonate releases gas by mposition.

$$NH_4HCO_3 \rightarrow NH_3 + H_2O + CO$$
,

are used as a leavening agent by bakeries for cookies and are occarily added to cream puffs and eclairs to improve their volume. The amplia formed adds a faint flavor and raises the pH so that the color and ading of the cookie are affected. In home baking ammonium bicarboor carbonate is seldom used because it decomposes readily and is difficultion of ammonium carbonate. Since ammonium bicarbonate reacts kly, it is sometimes added to produce rapid leavening. In preparing cookies which are topped with marshmallow, a soft, well leavened, or thick cookie is required. In the oven the ammonium bicarbonate a quick spring before the cookie spreads excessively.

r is a most important leavening agent in many types of batters. In a cake and angel cake it is traditionally but not always actually the leavening agent. The old pound cake recipe which used a pound each otter, sugar, flour, and eggs is leavened by thorough creaming of the ad sugar. As the mix becomes fluffy, small bubbles of air are adsorbed in the oven the expansion of these bubbles raises the cake. Often today amounts of other leavening agents are added in order to increase the me and lightness of the cake. In angel cake the egg white foam holds ir, and mixing is controlled so that the maximum amount of air is intorated and the minimum lost while the other ingredients are added. In ge and butter cakes baking powders are used to produce carbon die and achieve leavening, but air is also important and the amount intorated in the egg white or whole egg influences the final volume of the

cream puffs and eclairs occasionally have small amounts of other ning agents added to increase the volume produced. Steam is import in all batters since the vapor pressure of water increases rapidly increased temperature. During the cooking of all batter products, her doughnuts, griddle cakes, or oven heated cakes, the steam formed to the agents that gives a light porous structure.

Retention in Batters. Batters are almost always extremely complex res and the specific effect of many components of these systems is for nost part only partially understood at present. The production of a nost part of partially understood at present, the attainment tender, high volume cake, doughnut or muffin requires the attainment delicate balance between elasticity and rigidity, cohesiveness and

crumbliness, tenderness and toughness. Recipes for batters have develor through empirical methods where the cook tried various combinations stumbled on some that were acceptable. Today enough research has go into batter products so that some rules of thumb for balancing conformulas, for instance, have been put together. Although the effect of exing edient is known in a general way, the day when these effects can be plained by physical chemistry is still some way off. Those factors when we have the batter elastic so that it can stretch around gas bubbles and allow them to coalesce or escape from the top, are: (1) the gluten developing the flour, and (2) the proteins of milk and eggs, particularly expressed that decrease elasticity and consequently prevent a rubbery textual thus increasing tenderness, are: (1) fats and (2) sugar.

Flour is the source of the gluten in the batter and consequently the ty of flour and all its characteristics has a most important bearing on qualities of the batter product. In bread flours a high quality gluten is sirable, but in batter products the amount of gluten development possi and the quality should be low. In general, flours produced from soft why ield better cakes than those from hard wheats. The gluten from swheats is much weaker than that from hard wheats and often, but always, the total quantity of protein is lower. Flour formed in the short separation during milling, which contains the highest amount of star and the lowest amount of protein, yields the best cakes. The granulation the flour also has considerable effect. The finer the granulation, the high the score of cakes baked with it. Cake flour is as finely granulated as economically feasible.

The amount of mixing has an effect on the extent of gluten development and consequently is one of the factors which contributes to the strength the batter. Lowe¹³ and her students have found that with a standard carecipe cake volume increases with mixing up to a maximum and then coreases. The batter of the overmixed cakes flows from the spoon in logithesis. The batter of the overmixed cakes flows from the spoon in logithesis, suggesting that gluten has been well developed. Methods of ming, whether creaming, blending, or muffin method, yield cakes with different volumes and different scores (see p. 347).

The effect of egg protein on the structure is well demonstrated who muffins or one-egg cake is compared with cake made by convention recipes. The grain tends to improve with the addition of egg up to a maximum and then, although the grain remains fine, the volume decreases at the texture becomes rubbery. It is also possible that the milk proteins a important in the final structure. The gluten and other proteins have the effects here—elasticity in the batter and rigidity after baking. In the bat it is essential that some elasticity occur to allow formation, retention as

rsion of gas bubbles. It is probably the gluten which develops during ng that is primarily responsible for elasticity in the batter. Although ability of egg white and of whole egg to form elastic films is demond when egg-white foams or whole-egg foams are prepared, the tenactives foams is not great. This can be observed when the foam is interested into the batter. Overmixing and rough handling must be ded so that loss of gas does not occur. During baking the proteins are tured and a rigid structure results which maintains itself around the pubbles even after the cake has cooled and the gas in the bubbles has racted. Milk and egg proteins probably augment the effect of the flour eins.

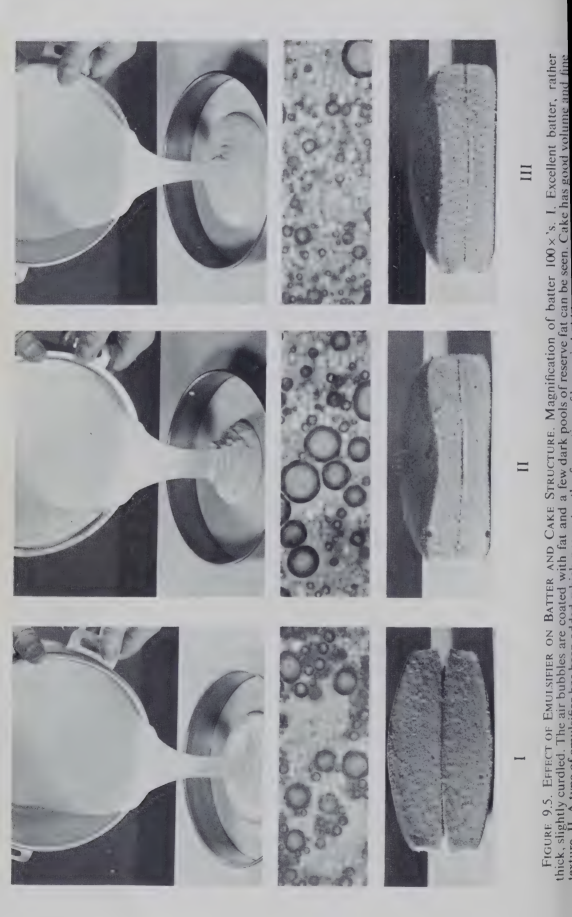
y fats available today, particularly the hydrogenated fats, contain sifying agents. These agents not only emulsify the fat so that it is redistributed through the batter, but probably also function to help not the tiny bubbles of gas in the batter. Moreover the fat interferes the development of gluten. This allows the protein particles to slide on another and the product is tender when bitten or cut.

o- and diglycerides as well as other esters of fatty acids such as sorbityl ate and the lecithins are useful. The addition of these emulsifying ts at 3 per cent of the total fat causes an increase in the volume of the as well as in the overall quality score. These agents also permit the tion of larger amounts of sugar and of moisture to the mixture, prong a sweeter, moister cake which is softer, which can retain moisture er, and has better keeping qualities. Batters that are made with fats aining emulsifier do not show a tendency to separate or break into

udy of the structure of the cake by staining in order to identify fat, h, or protein shows that the fat tends to occur in tiny lakes surround-he gas bubbles with some distribution through the crumb. The method ixing, the temperature, and the kind of fat influence the distribution of

ough the cake.

gar. In cakes and cookies sucrose is one of the principal ingredients h makes for tenderness. In the presence of sucrose development of n s retarded and restricted. Cakes in which the weight of sucrose is to or exceeds that of the flour (by measure, 1 2 cup sugar to one cup are called high-sugar cakes and are very popular. Those in which the nt of sucrose is less than that of flour are called low-sugar cakes. High-reakes are very sweet and have a fine texture with few tunnels. Cakes high both the sugar and moisture are high (moisture includes the liquid



texture II A time of emulcifier has been added within

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ed in milk and eggs) are high-ratio cakes and are possible today because the emulsifying agents added to fats. (See p. 342) Lowe and her sturing than low-sugar cakes in order to achieve optimum volume during and prevent falling when removed from the oven. Too much sugar as a sticky crust and a gummy texture.

igar has a marked effect on the hydration of proteins and starch which are in any batter or dough and must through this competition for water if the physical properties of the baked product. Proteins hold water on surfaces and within the folded or coiled structure of the molecule by ogen bonding. A well-hydrated protein is probably a tender one in the adherence of one protein molecule to another is weak and easily lered. The starch hydrates and forms a gel in doughs and batters, pararly during baking. The extent of gel formation must also have a ked effect on the physical properties of the baked goods. Sucrose as all molecule with many polar groups has an attraction for water and compete with protein and starch molecules for it. Undoubtedly the rences in high-sugar and low-sugar cakes reflect the competition of ar, protein, and starch molecules for water.

1. The effect of pH on cake quality has been studied and the effect on are observed, but there has been no direct study of the influence of pH



T SUGAR BUT WITH SU-It is difficult to produce grain and volume without The recipe must be very ly balanced.

rie v of Abbott Laboratories.

on gas retention. Stamberg and Bailey²⁰ reported that the optimum plant for white and yellow cakes could be narrowed to 7.22–7.35. These cakes were made with either phosphate or tartrate baking powders and those is which the pH was markedly different were inferior.

The ingredients which influence the pH of a white or yellow cake are the baking powder and the buffer salts which are present in milk. Disodiur phosphate formed from a phosphate baking powder will give a pH a little above 7, depending on its concentration. Calcium phosphate is quite in soluble and has little effect. Sodium potassium tartrate from tartrate bak ing powders gives a pH above 7. Sodium sulfate, the product of an S.A.S baking powder, gives a pH very close to 7. If bicarbonate of soda is adde to any recipe, it raises the pH and if sodium carbonate is formed durin baking from any soda which has not reacted with acid the pH is high. Acid are added to some cakes (but not usually to white or yellow) by use of sou milk, molasses, honey, or chocolate and cocoa. Since the amount of acid varies with the sample, it is very difficult to use these and achieve exact o close to exact neutralization of the soda. If the acids are present in insuffi cient amounts, the cake will be alkaline and have a high pH; if they are in excess, part of these acids will vaporize during baking but the pH of the cake will be lower than if they had not been added. Cream of tartar (potassium acid tartrate) is sometimes added to lower pH. Daniels and Heisig measured the acidity of some of the syrups used in baking, many years ago and found considerable variation. One cup of molasses varied from 12.8 to 22.6 ml of 0.1 N NaOH for neutralization.

Stamberg and Bailey²⁰ give the following pH values for commercial cakes:

White Cake	7.47
Yellow	7.59
Chocolate	8.48*
Angel Food	5.67

^{*}Must have been made with soda.

Cathcart⁴ gives the following pH ranges for commercial cakes:

Angel Food	5.2-5.6
Chocolate	7.2-7.6?
Devil's Food	7.5-8.0
Fruit Cake	4.4 - 5.0
Pound	6.6-7.1
Sponge	7.3-7.6
White	7.1-7.4
Yellow	6.7 - 7.1

re texture of a cake changes with changes in pH, tending to be finer and compact at lower pH while it becomes coarser on the alkaline side would expect from our knowledge of the change in characteristics of ns with change in pH that the elasticity and toughness of the proteins four, milk, and eggs are influenced by pH. This variation of texture pH must surely reflect effects on protein.

general the flavor and overall quality score for cakes is better when H is lower. The length of time for the development of staling is like-

prolonged with lower pH.

ocolate and Devil's Food cakes have different meanings to different rs and a wide range of pH values are found for commercial as well as e made cakes. (Hence the question mark which Cathcart has placed the pH range given for chocolate cakes.) As the pH rises, the color ges through cinnamon brown to brown and then becomes more and red. The flavor of the chocolate likewise changes. Many bakers add uch extra soda to Devil's Food cake that the pH is above 8, the color ry red, and the flavor very poor. The bitter soapy taste of sodium onate is readily detected and is most objectionable.

king Temperature and Time. Numerous studies have been made of the ; of baking temperature and time on the qualities of cakes. It is imant that the batter temperature rise so that baking powder, particularly 5. powders which react very slowly at room temperature, will form and that the gases will expand. It is also essential that the walls set. at the gas is not lost and the bubbles do not coalesce and form a coarse

In general it has been found that a relatively high temperature and equently a short baking time starting with a preheated oven give the 's' volume and the highest scoring product. Charley' found that a cake In Japanned iron had a better volume when baked 24 minutes at F than 27 minutes at 345° F. In tinned iron she found a temperature of F preferable.

visture. The amount of moisture in a cake and the manner of its distion are doubtless factors important to the quality of a batter product. eeling in the mouth of moistness or of dryness adds a great contribuo the enjoyment of eating. Moisture is held in a batter or dough prodthe solvent for some compounds, particularly sugar, and as a hydrate of an and starch. During baking, proteins are denatured and the in of water held in the molecule changes. The denatured protein can ld water through hydrogen bonding. (See p. 122.) Starch swells and atmized. The granules swell, water passes into the molecules, and is here (see p. 104).

Staling

Staling is a process which occurs in all dough and batter products renders them less desirable for eating. Attempts to understand and co the process have been made for many years; and although some pro has been made, there is still a lot to learn. Most of the work has bee bread, but the same process occurs in any flour mixture. In bread the loses its crispness and becomes leathery, while the crumb becomes and crumbly and dries out during the first two or three days after ba The hard crumb first appears just under the crust, but with loss of moi it progresses through the whole loaf. At the end of two or three week entire loaf has become hard and dry. Early in the process the staling ca reversed by heating the loaf. Quantitative estimates of staling have devised using (1) a penetrometer which gives a measure of the hardne the crumb and (2) the amount of swelling which occurs when a crun placed in water. Fresh bread swells much more extensively than stale. bread which is stored at 60°C or at very low temperature staling is much delayed. But the flavor at 60°C suffers deterioration. Freezing b and storing it well below the freezing point is a practical and much method of maintaining baked products for considerable periods of t The most rapid staling occurs at -2° C to -3° C.

Staling appears to be associated with changes in the starch. Chemic there is a loss in the amount of extractable starch which is obtained to bread as staling proceeds. Some workers believe that it is the relatively molecular weight amylopectins which are involved while others believe the straight chain amyloses. Although the process is not completely ar gous to the changes which occur when a simple starch-water paste of undergoes retrogradation, there are many indications that the proce similar. Today most investigators conclude that the basis of similarity tween staling and retrogradation is that both are caused by a reorientate of the hydrogen bonds. It is thought that during both processes water in cules associated with starch molecules are lost. This leads in retrogration to marked orientation of the starch molecules and the development crystalinity. In staling the orientation does not appear to be as complete.

The addition of emulsifying agents such as mono- and diglycerides other compounds similar to them lengthens the keeping quality of b and delays staling. They are now extensively used in the United States. up to 2 per cent of the flour improve bread and slightly retard the sta of the loaf. Addition of protein to the dough gives a softer crumb b stronger structure to the loaf and produces a slight delay in staling. Pa the delay is apparent because of the initial softness of the crumb.

taling retardants are appearing in the patent literature. In 1956 U.S. ent 2,744,825 was issued to Thompson and Buddenmeyer for fatty acid ylates. These compounds are patented for use in yeast-leavened doughs incentration of 0.1 to 1 per cent of the weight of the flour. Their gentructure is

ere RCO is a fatty acid chain with 16 22 carbon atoms, n varies from 8 and X is a cation.

oring Baked Products

Many scoring forms have been developed by industries producing large ounts of baked products, by laboratories, and others interested in luating quality. Sometimes a rating of 1 to 10 is used for each attribute, I sometimes other numbers. Whatever the method a comparative nurical score is obtained that is useful in controlling the quality of the ducts or evaluating changes in methods and ingredients. Usually the ilities evaluated are: appearance, color of crust and crumb, aroma, ture, moistness, tenderness, flavor, symmetry and general eating quality. casionally odor and flavor may be combined since they are so interned and often reflect the same quality. Appearance is sometimes divided general appearance including smoothness, greasiness, pebbling and 'ary as well as the general contours such as humped, sunk, flat, etc. and another evaluation for the volume of the product such as too large. od, too small. Texture refers to the grain, the thickness or thinness of cell walls, the size of the gas bubbles, and occasionally includes a asure of tenderness.

evention of Mold

Prevention of mold on bakery products has been the object of much rerch. Mold growth, which causes considerable loss, develops in wrapped ods when humidity is high and temperature also fairly high. The baked do are sterile when they leave the oven; but as they cool and are sliced wrapped, they are contaminated by mold spores from the air. Air containing which brings in washed air and meticulous cleanliness in the cooland wrapping rooms reduce the number of mold spores in the air and refore the amount of contamination. Even so, some molding will occur.



FIGURE 9.7. MOLD GROWTH ON BREAD.

Courtesy of Fleischmann Laboratories. Standard Brands

Numerous compounds can be added to the dough or batter in enough concentration so that toxicity or flavor will not occur and you high enough concentration so that mold growth is sharply inhibited. dium and calcium propionate are now widely used in the baking indu

CH₃CH₂COONa Sodium Propionate CH₃CH=CHCH=CHCOOH
Sorbic Acid



FIGURE 9.8. MOLD GROWTH ON BI 'Magnification × 25. Courtesy of Fleischmann Laborat Standard Brands

TABLE 9.4. MOLD INHIBITION

Suppleme	nt to the Batter		Mold Gr after I	Mold Growth in Humidity Can after Days Indicated 90° F			
unce	Concentration %	2	4	5	7		
2	0.00	_	++++	++++	++++	++++	
ic Acid	0.05	_	++	+++	++++	++++	
	0.10	_	_	_	+	+++	
	0.20		_	_		_	
ropionate	0.30		+	++	+++	++++	
	0.60	_	_	+	++	++++	
senzoate	0.1	_	++	++++	+++-	+ + + +	

om Melnick, D., Vahlteich, H. W., and Hackett, A., "Sorbic Acid as a Fungistatic Agent for Loods," Research, 21, 133-146 (1956).

chieve this purpose. In more recent years sorbic acid and its salts have studied and appear to be at least as effective as the propionates. nick, Vahlteich, and Hackett14 added mixed mold culture to a variety ake batters and measured the effectiveness of some compounds as intors of mold growth in slices of cake stored in warm humid containers. r results with white cake are shown in Table 9.4 and indicate good conat low concentration.

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JAPTER TEN

ood Additives

MOST FREQUENTLY DISCUSSED food chemistry problem in the popular stoday is that of food additives. A food additive is any substance not ally present in a food but added during its preparation and remaining the finished product; also in this category is any substance naturally sent but with a concentration increased by fortification. However since sugar, and vinegar have been used for centuries, they are not usually sidered food additives. Today many compounds are added to foods and 1st of substances that protect against spoilage, that enhance flavor, that prove nutritive value, or that give some new property to a food is insing rapidly.

he market for these food additives is large and growing. Since in the ited States the distance between producer and consumer of food can be at, the length of time between harvest and eating can present many biems in maintaining freshness. The economic status of the average content has improved until he is able to demand and pay for high grade ducts. There is a constant search for methods to improve quality: ways, example, to prevent a fall-off in flavor as poultry or fish stands in the cher's cold counter; to prevent the high loss of fresh fruits and veges between field and kitchen; and to prolong the shelf life of bakery ducts. The list of similar problems could be extended at length. Added his is the highly competitive nature of the food industry where yesteris methods and products do not necessarily mean success in today's act.

FAL SAFEGUARDS

is the production of new food additives increased, suspicion grew that guards against toxicity were not adequate. Considerable interest de-

veloped in the methods used by federal agencies to ensure the safet additives. The toxicity of pesticides for insects alarmed some cowho believed that surely these compounds that kill insects might be lethal for themselves.

Agene, nitrogen trichloride, was used as a bleach for flour for years. Bleaching improves the baking quality of flour in a short the produces a flour comparable to the more expensive aged product was used in very low concentration but it was shown in the late a flour bleached with this agent or bread baked from this flour cause ning fits in dogs and convulsions in rabbits, ferrets, mink, and cat port which led to an almost hysterical distrust of all food additionall segment of our population. The animals adversely affected which diets composed principally of the flour or bread. The toxic computation reproduced from the action of nitrogen trichloride on methionine methionine sulfoxamine. It has never been demonstrated that harr fects are caused by this flour or bread in man although a number of ments have been conducted to test the possible toxicity. Neverthelective August 1, 1949, agene was removed from the list of periodical periodical agents.

After much debate and a number of hearings and bills, the Amendment was passed in 1954 and the Food Additives Amendmen lic Law 85–929) in 1958. The Miller Amendment covers spray residuthe use of pesticides; it attempts to establish procedures that will any possible harm to consumers. The Food Additives Amendment requirements for the demonstration of the safety of a substance it can be used in foods; it went into effect on March 5, 1960, althoug items have had an extension beyond this date.

The Food Additives Amendment uses the following definition:—additives' include all substances not exempted by section 201 of the intended use of which results or may reasonably be expected to directly or indirectly, either in their becoming a component of fotherwise affecting the characteristics of food." The container is n sidered a food additive but if some of the substances present in that tainer migrate into the food, then they become food additives and to the law.

A list of additives that are already in use and appear to have been quately shown to be harmless has been prepared. No further testing quired with these compounds that are considered permissible. The Appendix, p. 357) includes preservatives, buffers and neutralizers, ents, non-nutrient sweetners, coloring agents, stabilizers, emulsified others. Although salt, sugar, and vinegar are not included, they are

ted. A list of permissible flavor substances is in preparation. Other adves cannot be used unless permission is granted by the Food and Drug ninistration. A petition for use must include a thorough review of the rance, its stability under all conditions of possible use, and a complete of the additive or its degradation products if they are formed in the J must also be given. The Food and Drug Administration guarantees ection of trade secrets. Only those facts about the substance that are irred for its control will be revealed.

IY USE FOOD ADDITIVES?

t a meeting of the Institute of Food Technologists in May, 1958, old Schultz6 is reported to have said of food additives that "some are to color foods, others to bleach them; some add flavor to food, others ove flavors; some make foods firmer, others soften them; some keep ! dry, others keep them moist; some thicken foods, others keep them n thickening; some produce foams, others prevent them; some make is acid, others make them alkaline, still others suspend particles; some .dded to increase the mineral ions, others to remove them; some are tizing agents, others are reducing agents; some hasten chemical nges, others retard them." This is a formidable list of functions aligh sometimes one food additive has more than one. It is a list that ilrates the needs of the food industry in its efforts to produce foods with cient shelf life to allow distribution over a wide area and with sufficient eal to the eye and taste of the consumer to arouse and satisfy his apte. These food products must nourish the eater well; but before they nourish him, they must first attract his purchase. Justifiable uses of ditives have been described in some detail by the Food Protection nmittee in Publication 398, "The Use of Chemical Additives in Food cessing." They are:

) "maintenance of nutritional quality, such as by the use of antioxidants."

"enhancement of keeping quality or stability, with resulting reduction in food wastage, through the use of antioxidants, antimicrobial gents, inert gases, meat cures, etc."

"enhancement of attractiveness of foods by means of coloring and flavoring agents, emulsifiers, stabilizers, thickeners, clarifiers, and bleaching agents. (That appeal to the eye as well as the palate is not a mere whim of the food producer, is recognized by all nutrition-

- ists, since proteins, vitamins, and minerals serve no purpo eaten.)"
- (4) "providing essential aids in food processing. Agents which in this capacity include acids, alkalis, buffers, sequestra various other types of chemicals."

METHODS OF DEMONSTRATING SAFETY

The current methods of demonstrating the safety of a chemic pound or crude extract to be used in food production are fa standardized. Those compounds which are not normally present foods are fed in graded amounts to rats for two years and to one year. The control animals are fed complete diets devoid of ditive. The level of feeding at which injury can be demonstrate termined.

The success or failure of this method depends on recognition jury." A considerable amount of attention has been devoted to thi The Food and Drug Administration uses the distinction between and hazard set forth by the National Research Council's Food a trition Board: "Toxicity is the capacity of the substance to produc hazard is the probability that injury will result from the use of stance in the quantity and in the manner proposed." Since evaluinjury depends on subjective judgment when changes are slight, trequires a complete description of methods used. "No effect" supplemented by a description of the conditions under which thereffect." The decision of the safety, toxicity, or hazard of a chemical distribution of the safety and the safety of the safety of the safety.

For some substances it is possible that use in small amounts probut in very large amounts hazard exists. Even a substance such as chloride, commonly considered completely safe to add to foods, compounds a limit is established by adding a margin of safety to the level. This margin must include consideration of the difference between man and test animals, individual variation, cumulative effects the possibility of the compound occurring in other items in the diagain judgments rather than measurements are used and much opossible.

It is unlikely that most chemical additives will be chemically pupounds. FDA requires that methods for the detection and estiming impurities be developed and presented with the petition for perm

Although a pure substance might be safe, the impurities carried along ht be injurious.

here is also a possibility that a compound might be sale as such but a such processing of a food, it could cause a change in some coment to form a poisonous compound. The toxic factor in flour bleached agene proved to be a derivative of methionine rather than the agene it is therefore required that a compound not only be demonstrated but also safe under the conditions of its use. Permissible chemical trives are restricted to those particular foods or processes which meet se requirements.

LOR ADDITIVES LEGISLATION

the FDA is requesting legislation to modify the present regulations red ng color additives to foods (as well as to drugs and cosmetics.) At sent color additives are either classed as "harmless" or not, with no red for the quantities used. Since this is not very realistic, the FDA hes controls set up so that the quantities in specific foods can be limit to safe levels. They also wish the law extended to include not only syntic coloring substances but also natural pigments used as additives.

Regulations on food additives and explanations of the law and of atpts to put it into smooth operation are being published by the FDA. ese publications are available free of charge to anyone who asks that name be placed on the mailing list.

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ppendix on ood Additives*

g Administration is now issuing lists of permitted food additives. The wing lists include those published in the Federal Register on Novem-20, 1959, and lists of January 19 and 26, 1960. It is expected that the plant Drug Administration will continue to issue lists of permissible jadditives.

TAIN FOOD ADDITIVES EXEMPTED FROM REQUIREMENT OF TOLERANCES

stances That Are Generally Recognized As Safe

It is impracticable to list all substances that are generally recognized are for their intended use. However, by way of illustration, the Commistregards such common food ingredients as salt, pepper, sugar, vine-biking powder, and monosodium glutamate as safe for their intended. The lists in paragraph (d) of this section include additional substances when used for the purposes indicated, in accordance with good manuring practice, are regarded by the Commissioner as generally recognas safe for such uses.

For the purposes of this section, good manufacturing practice shall

ef ned to include the following restrictions:

The quantity of a substance added to food does not exceed the art reasonably required to accomplish its intended physical, nutri-

al, or other technical effect in food; and

The quantity of a substance that becomes a component of food as a tefits use in the manufacturing, processing, or packaging of food, and bis not intended to accomplish any physical or other technical effect in

ood itself, shall be reduced to the extent reasonably possible.

The substance is of appropriate food grade and is prepared and as a food ingredient. Upon request the Commissioner will offer an ased on specifications and intended use, as to whether or not a lar grade or lot of the substance is of suitable purity for use in food a suld generally be regarded as safe for the purpose intended, by exqualified to evaluate its safety.

^{*} the Federal Register, U. S. Department of Health, Education, and Welfare, Food Administration, Excerpts of November, 1959 and January, 1960

- (c) The inclusion of substances in the list of nutrients does not a finding on the part of the Department that the substance is us supplement to the diet for humans.
- (d) Substances that are generally recognized as safe for their use within the meaning of section 409 of the act are as follows:

Chemical Preservatives

Ascorbic acid. Ascorbyl palmitate. Calcium ascorbate. Calcium propionate. Calcium sorbate. Erythorbic acid. Potassium sorbate. Propionic acid. Sodium ascorbate. Sodium propionate. Sodium sorbate. Sorbic acid. Tocopherols.

Buffers and Neutralizing Agents

Acetic acid.

Adipic acid.

Aluminum ammonium sulfate.

Aluminum sodium sulfate.

Aluminum potassium sulfate.

Ammonium bicarbonate.

Ammonium carbonate.

Ammonium hydroxide.

Ammonium phosphate (mono- and di-

Calcium carbonate.

Calcium chloride.

Calcium citrate.

Calcium gluconate. Calcium hydroxide.

Calcium lactate.

Calcium oxide.

Calcium phosphate.

Citric acid.

Hydrochloric acid.

Lactic acid.

Magnesium carbonate.

Magnesium oxide.

Malic acid.

Phosphoric acid.

Potassium acid tartrate.

Potassium bicarbonate.

Potassium carbonate.

Potassium citrate.

Potassium hydroxide.

Sodium acetate.

Sodium acid pyrophosphate.

Sodium aluminum phosphate.

Sodium bicarbonate.

Sodium carbonate.

Sodium citrate.

Sodium hydroxide.

Sodium phosphate (mono-, di-, tr

Sodium potassium tartrate. Sodium sesquicarbonate.

Succinic acid.

Sulfuric acid.

Tartaric acid.

Emulsifying Agents

Diacetyl tartaric acid esters of diglycerides from the glycerolys fats and oils.

Mono- and diglycerides from the

sis of edible fats or oils.

Monosodium phosphate deri mono- and diglycerides from th sis of edible fats or oils.

Propylene glycol.

Miscellaneous

Acetic acid.

Aluminum sodium sulfate.

Aluminum sulfate.

Ammonium bitartrate.

Ammonium sulfate. Bees wax (yellow wax).

Bees wax bleached (white wax).

Bentonite.

Butane.

Calcium chloride.

Calcium phosphate, tribasic.

Caramel

Carbon dioxide.

Carnauba wax.

Citric acid. Dextrans (of average molecular

low 100,000).

Glycerin.

Glycerol monostearate.

Helium.

Lecithin.

Magnesium carbonate.

Magnesium hydroxide.

Methyl cellulose (U.S.P. methylce cept that the methoxy content s less than 27.5 per cent and not 31.5 per cent on a dry-weight ba vammonium glutamate. notassium glutamate.

wen.

roric acid. ...um carbonate. ssum sulfate.

wene glycol. et (rennin).

im bicarbonate. m carbonate.

im carboxymethylcellulose (the sodium of carboxymethylcellulose not less ... 99.5 per cent on a dry-weight basis, th maximum substitution of 0.95 carbymethyl groups per anhydroglucose 1 and with a minimum viscosity of centipoises for 2 per cent by weight reous solution at 25° C.)

.m caseinate. um phosphate.

um sesquicarbonate.

pectinate. am phosphate.

m tripolyphosphate. na yeast.

e in (Glyceryl triacetate). . kium phosphate.

nutritive Sweeteners

nonium Saccharin rum cyclohexyl sulfamate. um saccharin. nesium cyclohexyl sulfamate.

sium cyclohexyl sulfamate.

um cyclohexyl sulfamate. am saccharin.

this acid.

ients

um carbonate.

sum oxide.

im pantothenate.

um phosphate (mono-, di-, tribasic).

" sulfate.

* 449 .0

ne bitartrate. phosphate.

- p rephosphate.

c sodium pyrophosphate.

sulfate.

reduced.

monohydrochloride.

ha nide.

ntothenyl alcohol.

in chloride.

mine hydrochloride.

Riboflavin.

Riboflavin-5-phosphate

Sodium pantothenate

Sodium phosphate (mono-, di-, tribasic)

Thiamine hydrochloride. Thiamine mononitrate.

Tocopherols.

α-Tocopherol acetate.

Vitamin A.

Vitamin A acetate.

Vitamin A palmitate.

Vitamin B₁₂.

Vitamin D₂.

Vitamin D₃.

Sequestrants

(For the purpose of this list, no attempt has been made to designate those sequestrants which may also function as chemical preservatives)

Calcium acetate.

Calcium chloride.

Calcium citrate.

Calcium diacetate.

Calcium gluconate.

Calcium hexametaphosphate.

Calcium phytate.

Citric acid.

Dipotassium phosphate.

Disodium phosphate.

Monocalcium acid phosphate.

Monoisopropyl citrate.

Potassium citrate.

Sodium acid phosphate.

Sodium citrate.

Sodium diacetate.

Sodium gluconate.

Sodium hexametaphosphate.

Sodium metaphosphate.

Sodium phosphate (mono-, di-, tribasic-).

Sodium potassium tartrate.

Sodium pyrophosphate.

Sodium tartrate.

Sodium tetrapyrophosphate.

Sodium tripolyphosphate.

Tartaric acid.

Stabilizers

Agar-agar.

Acacia (gum Arabic).

Ammonium alginate.

Calcium alginate.

Carob bean gum (locust bean gum).

Carragheenin.

Ghatti gum.

Potassium alginate.

Sodium alginate.

Sterculia gum (Karaya gum).

Tragacanth (gum tragacanth).

Guar gum.

Substances Allowed If Restrictions Are Observed

Product	Tolerance	Specific uses or restrict
ANTICAKING AGENTS		
Aluminum calcium silicate	2 per cent	In table salt. In baking powder.
Calcium silicate	2 per cent	In table salt. Do. Do.
CHEMICAL PRESERVATIVES		
Benzoic acid	0.1 per cent	
Butylated hydroxytoluene	do	In cheese wraps.
Dilauryl thiodipropionate	Total content of antioxidants not over 0.02 per cent of fat or oil content, including essential (volatile) oil content of the food.	meneese wraps.
Gum guaiac	0.1 per cent (equivalent antioxidant activity 0.01 per cent).	In edible fats or oils.
Nordihydroguaiaretic acid	Total content of antioxidants not over 0.02 per cent of fat or oil content, including essential (volatile) oil content of the food.	
Potassium bisulfite	• • • • • • • • • • • • • • • • • • • •	Not in meats or in food re as a source of vitamin B ₁
Potassium metabisulfite Propyl gallate	Total content of antioxidants not over 0.02 per cent of fat or oil content, including essential (volatile) oil content of the food.	Do.
Sodium benzoate	0.1 per cent	Not in meats or in foods re
Sodium metabisulfite		as a source of vitamin B_1 .
Sodium sulfite		Do.
Sulfur dioxide	Total content of antioxidants not over 0.02 per cent of fat or oil content, including essential (volatile) oil content of the food.	Do.
EMULSIFYING AGENTS		
Cholic acid	0.1 per cent	Dried egg whites.
Deoxycholic acid	do	Do. Do.
Ox bile extract	do	Do.
Taurocholic acid (or its sodium salt) . MISCELLANEOUS	do	Do.
	0.02	T 1 . 1
Caffeine, Ethyl formate	0.02 per cent	In cola type beverages. As fumigant for cashew nuts
Magnesium stearate		As migratory substance fr aging materials when used bilizer.
Sorbitol	7.0 per cent	In foods for special dietary u Egg whites.
NUTRIENTS		,
Copper gluconate	0.005 per cent	In table salt as a source iodine.
Potassium iodide	do	Do.
	0.02	
Isopropyl citrateSodium thiosulfateStaryl citrateStaryl citrate	0.15 per cent	In salt.

¹For the purpose of this list no attempt has been made to designate those sequestrants which may also function cal preservatives.

CES, SEASONINGS, ESSENTIAL OILS; OLEORESINS, AND NATURAL TRACTIVES THAT ARE GENERALLY RECOGNIZED AS FE FOR THEIR INTENDED USE.

ces and Other Natural Seasonings and avorings (Leaves, Roots, Barks, Berries, etc.)

Common name

Botanical name of plant source

pice	Pimenta officinalis Lindl.
SC	Pimpinella anisum L.
se, star	Illicium verum Hook. f.
il, sweet	Ocimum basilicum L.
il, bush	Ocimum minimum L.
	Laurus nobilis L.
	Calendula officinalis L.
	Capparis spinosa L.
sicum	Capsicum frutescens L. or Capsicum annuum L.
away	Carum carvi L.
way, black (black cumin)	Nigella sativa L.
damom (cardamon)	Elettaria cardamomum Maton.
sia, Chinese	Cinnamomum cassia Blume.
sia Padang or Batavia	Cinnamomum burmanni Blume.
s a. Saigon	Cinnamomum loureirii Nees.
enne pepper	Capsicum frutescens L. or Capsicum an-
	nuum L.
ry seed	Apium graveolens L.
ie	Allium schoenoprasum L.
namon, Ceylon	Cinnamomum zeylanicum Nees.
ramon, Chinese	Cinnamomum cassia Blume.
namon, Saigon	Cinnamomum loureirii Nees.
ry (clary sage)	Salvia sclarea L.
ves	Eugenia caryophyllata Thunb.
ander	Coriandrum sativum L.
in (cummin)	Cuminum cyminum L.
nin, black (black caraway)	Nigella sativa L.
.,	Anethum graveolens L.
nel common	Foeniculum vulgare Mill.
nel sweet (finocchio, Florence fennel)	Foeniculum vulgare Mill. var. dulce (DC.)
Mer. Sweet (Milocomo, 1 101011)	AIEI
uş-eek	Trigonella foenum-graecum L.
IIC	Allium sativum L.
ger	Zingiber cincinale Rosc.
yrrhiza	(IIVC VII IIII Zu Bisson
.ylilli.a	Glycyrrh173
ins of paradise	Amomum melegueta Rosc.
ser idish	Armoracia lapatillolla Gino.
nder	
inder	Glycyrrhiza glabra L. and other spr
rice	Glycyrrhiza.
	Myristica fragrans Houtt.
e	Calandula officinalis L.
old. pot	Majorana onites (L.) Benth.
vam, sweet	Majorana horiensis Wicelian
and block or brown	Registica nigra (L.) Noch
ard, black or brown	Denocios junces (1) (USS.
.tard, brown	Demostop alba (I DOID).
taid, white or yellow	Myristica iragians rivers
meg	Lippia spp.
ano (oreganum, Mexican oregano,	
exican sage, origan).	
	261

Paprika	Capsicum annuum L.
Parsley	
Pepper, black	
Pepper, cayenne	Capsicum frutescens L. or Caps
11	nuum L.
Pepper, red	Do.
Pepper, white	Piper nigrum L.
Peppermint	
Poppy seed	
Pot marigold	
Pot marjoram	Majorana onites (L.) Benth.
Rosemary	
Rue	
Saffron	Crocus sativus L.
Sage	Salvia officinalis L.
Savory, summer	Satureia hortensis L. (Satureja).
Savory, winter	Satureia montana L. (Satureja).
Sesame	Sesamum indicum L.
Spearmint	Mentha spicata L.
Star anise	Illicium verum Hook. f.
Tarragon	Artemisia dracunculus L.
Thyme	Thymus vulgaris L.
Turmeric	Curcuma longa L.
Vanilla	
	J. W. Moore
Zedoary	Curcuma zedoaria Rosc.

Essential Oils, Oleoresins (Solvent-Free), and Natural Extractives (Including Distillates)

Common name	Botanical name of plant sour
Allspice	Pimenta officinalis Lindl. Prunus amygdalus Batsch, Prunus
•	L., or Prunus persica (L.) Batsch.
Angelica root Angelica seed	Angelica archangelica L.
Angelica stem	Do.
Angostura (cusparia bark)	Galipea officinalis Hancock.
Anise	Pimpinella anisum L. Ferula assa-foetida L. and relate Ferula.
Balsam of Peru	Myroxylon pereirae Klotzsch.
Basil	Ocimum basilicum L.
Bay leaves Bay (myrcia oil)	Laurus nobilis L. Pimenta racemosa (Mill.) J. W. Mo
Bitter almond (free from prussic acid)	Prunus amygdalus Batsch, Prunus L., or Prunus persica (L.) Batsch.
Bois de rose	Aniba rosaeodora Ducke.
Cananga	Cananga odorata Hook, f. and Tho
Capsicum	Capsicum frutescens L. and Capanuum L.
Caraway	Carum carvi L.
Cardamon seed (cardamon)	Elettaria cardamomum Maton. Ceratonia siliqua L.
Cascarilla bark	Croton eluteria Benn.
Cassia bark, Chinese	Cinnamomum cassia Blume.
Cassia bark, Padang or Batavia	Cinnamomum burmanni Blume.
Cassia bark, Saigon	
Celery seed	Apium graveolens L. Matricaria chamomilla L.

mile flowers, Roman or English (cam-	Anthamic and the
	Anthemis nobilis L
and bark	Prunus serotina Ehrh.
	Chicorium intybus I
on bark. Ceylon	Cinnamomum zevlanicium Nome
on bark, Chinese	Cinnamomum cassia Blume
non bark, Saigon	Cinnamomum loureirii Nees
man leaf, Ceylon	Cinnamomum zevlanicum Nees.
non leaf, Chinese	Cinnamomum cassia Blume
mon leaf, Saigon	Cumhanagan nardus P. Cymhanagan nardus P.
peels	Cymbopogon nardus Rendle. Citrus spp.
(cary sage)	Salvia sclarea L.
bud	Eugenia caryophyllata Thunb.
raf	Do.
tem	Eugenia caryophyllata Thunb.
decocainized)	Erythroxylum coca Lam. and other spp of
	Erythroxylum.
	Coffea spp.
rut	Cola acuminata Schott and Endl., and other
	spp. of Cola.
ıcer	Coriandrum sativum L.
r (cummin)	Cuminum cyminum L.
r.a bark	Galipea officinalis Hancock. Anethum graveolens L.
······································	
: le (esdragol, esdragon, tarragon)	Foeniculum vulgare Mill.
steek	Trigonella foenum-graecum L.
STOCK	Allium sativum L.
m. East Indian	Cymbopogon martini Stapf.
III.M. rose	Pelargonium graveolens L'Her.
	Zingiber officinale Rosc.
TN1/a	Glycyrrhiza glabra L. and other spp. of
	Glycyrrniza.
:fruit	Citrus paradisi.
	Psidium spp. Marrubium vulgare L.
cound	was to the classical state of the contract of
	- · · · · · · · · · · · · · · · · · · ·
າະ	Jasminum.
	T i and communic
er (herries)	Cola acuminata Schott and Endi., and other
าน"	spp. of Cola.
· le ives	Laurus mobilis L.
le*	Lavandilla Officilians Chart.
der, spike	Lavandula latifelia Vill. Hybrids between Lavandula officinalis Chaix Hybrids between Lavandula officinalis Chaix
da	and Lavandula latifolia Vill.
	allu Lavanidata
	Citrus Limon (L.) Burm. f. Cymbopogon citratus DC. and Cymbopogon
Frass	flexuosus Stapf.
	Chapterbig glabra L. and other spp of
c	(legerrh 173
	Citrus aurantifolia Swillgic.
	Caratonia Silialia 1.
	Marietica tragrams muutt.
in	
Fam. Sweet	a f ' a horiellala la locitoria
· · · · · · · · · · · · · · · · · · ·	Ilex paraguariensis of
rd	D-000100 SDH
71.	Citrus paradisi Macf.

Neroli, bigarade	Citrus aurantium L.
Nutmeg	Myristica fragrans Houtt.
Onion	Allium cepa L.
Orange, bitter, flowers	Citrus aurantium L.
Orange leaf	Citrus sinensis (L.) Osbeck.
Orange, bitter, peel	Citrus aurantium L.
Orange, sweet	
Origanum	Origanum spp.
Palmarosa	Cymbopogon martini Stapf.
Paprika	Capsicum annuum L.
Parsley	Petroselinum crispum (Mill.) Mansf.
Pepper, black	Piper nigrum L.
Pepper, white	Piper nigrum L.
Peppermint	Mentha piperita L.
Peruvian balsam	Myroxylon pereirae Klotzsch.
Petitgrain	Citrus aurantium L.
Petitgrain lemon	Citrus limon (L.) Burm. f. Citrus reticulata Blanco.
Petitgrain mandarin or tangerine	Pimenta officinalis Lindl.
Pimenta leef	
Pimenta leaf	Do. Chimaphila umbellata Nutt.
Pipsissewa leaves	Punica granatum L.
Prickly ash bark	Xanthoxylum (or Zanthoxylum) Am
THERE'S ASIL DALK	Mill. or Xanthoxylum clava-hercu
Rose absolute	Rosa alba L., Rosa centifolia
1030 40301410	damascena Mill., Rosa gallica L.,
	of these spp.
Rose (otto of roses, attar of roses)	Do.
Rose geranium	
Rosemary	
Rue	
Saffron	
Sage	
Sage, Spanish	Salvia lavandulaefolia Vahl.
St. John's bread	Ceratonia siliqua L.
Schinus molle	Schinus molle L.
Spanish sage	Salvia lavandulaefolia Vahl.
Spearmint	Mentha spicata L.
Spike lavender	Lavandula latifolia Vill.
Tangerine	
Tarragon	
Tea	Thea sinensis L.
Thyme	Thymus vulgaris L. and Thymus 2
771	gracilis Boiss.
Thyme, white	Do.
Tuberose	Polianthes tuberosa L.
Turmeric Vanilla	Curcuma longa L.
vaiiiia	Vanilla planifolia Andr. or Vanilla
Violet leaves absolute	J. W. Moore.
Violet leaves absolute	Viola odorata L.
Wild cherry bark Ylang-ylang	Prunus serotina Ehrh.
Zedoary bark	Cananga odorata Hook, f. and Thon Curcuma zedoaria Rosc.
Louding bark	Curcuma zedoaria Rosc.
Miscellaneous	
Wilder Mileous	
Соммон намо	Domination

Comi

Common name			Deri	ivation	
Civet (zibeth, zibet, zibetum)	Civet	cats,	Viverra	civetta	Schre
Cognac oil, white and green	Viv. Ethyl	erra zil	betha Sch thate, so-	reber.	

dex

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e bodies in milk, 298	Allylisothiocyanate, 279
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